EXHIBIT A

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MOLECULAR CRYSTALS AND LIQUID CRYSTALS

SUPPLEMENT SERIES.

SUPPLEMENT 1

ALIGNMENT OF NEMATIC LIQUID CRYSTALS AND THEIR MIXTURES

Jacques Cognard



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MOLECULAR CRYSTALS and LIQUID CRYSTALS

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Alignment of Nematic Liquid Crystals and Their Mixtures by Jacques Cognard

MOLECULAR CRYSTALS AND LIQUID CRYSTALS, SUPPLEMENT SERIES

SUPPLEMENT .

Alignment of Nematic Liquid Crystals and Their Mixtures

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Preface

Research in the field of Molecular and Liquid Crystals has reached a certain degree of maturity. It was considered timely, therefore, to inaugurate a supplement series to the journal of Molecular Crystals and Liquid Crystals. The aim of this series is to provide in-depth reviews, in the form of monographs, of specific topics within this wide and rapidly growing field. As a result of expansion and diversification, regular research articles must be highly specialized. It is important, therefore, to provide the mechanism whereby researchers, students and practitioners can readily obtain a balanced view of the important results in various areas of the overall field. This is the aim of this new supplement series.

G. J. Dienes Managing Editor

Alignment of Nematic Liquid Crystals and Their Mixtures

JACQUES COGNARD

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Tilted alignment
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INTRODUCTION

The most recent reviews concerning the alignment of liquid crystals (LC) appeared five years ago. 1-3 Since then studies of LC alignment have proceeded actively: the tangential evaporation of oxides and fluorides has found wide industrial acceptance: aligning properties of layers deposited in various ways have been well documented and studies of LC surface interactions are numer-

ALIGNMENT OF LIQUID CRYSTALS

ous. Many methods of aligning LCs have been described but attempts to utilize or even reproduce published results are often discouraging.

It is the purpose of this paper to review the various methods described, to extend them to other commercial LC mixtures and point out the most reliable ones, as found through our own experience in developing displays for electronic watches.

We have sought to be comprehensive up to December 1979 and cite any English, French or German references available to us through computer search, abstracts bibliography and cross referencing. It was thus impossible to cover all studies and we considered only the most practical methods of LC alignment compatible with practical requirements. We insist on those processes that give reproducible results.

An evaluation of an alignment process has to consider its reproducibility, applicability to various LCs, lifetime and compatibility with the associated

LC Code

(1) LC commercial mixtures

(from our chromatographic analysis)

Roche	105	0.31 M 15: 0.14 M 24; 0.14 T 15; 0.09 P ₃ 5 0.18 P ₃ 7: 0.14 T P ₃ 4	ROTN 404
Roche	65.2	0.33 53; 0.67 56	ROTN 200
Roche	81.4	0.05 PEPN 4: 0.24 PEPN5;0.13 PEPN 6 0.18 PEPN7: 0.12 P ₃ 5: 0.08 P ₃ 7: 0.20 T P ₃ 4	ROTN 103
врн	70.5	0.45K15; 0.16M9; 0.12M15; 0.16M24 0.11 T 15	€ 8
врн	59. 8	0.51 K15; 0.25 K 21; 0.16 M 24; 0.08 T 15	E 7
врн	54.3	0.55 K15; 0.14 M 15; 0.13 M 21; 0.18 M 24	E 3
Merck	70	0.15 BCH-5; 0.24 PCH-3; 0.36 PCH-5 0.25 PCH-7	Zli 1132
Source	T _{NI} °C	Composition	Code

Merck—Frankfurter Str. 250 D 6100 Darmstadt 1
BDH —Poole Dorset BH 124 NN England
Roche—Liquide Crystals group RA/LC CH-4002 Basel

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sealing process. Sealing of LC displays involves either a high temperature (500°C) glass frit technique or adhesives cured around 200°C.

Most alignment methods have been tested on MBBA,† a very sensitive Schiff's base LC, while biphenyls, esters or phenylcyclohexane eutectic mixtures are of current industrial interest. Our evaluations considered the commerical mixtures ROTN 200, ROTN 103, ROTN 404, E7, E8 and Zli 1132.† On filling a cell, the nematic director generally aligns in the direction of flow (4a), but the resulting orientation is temperature sensitive (4b). Heating the LC to its isotropic phase will often promote a uniform alignment, and slow

LC Code

(2) LCs of current industrial use

C 8 - 5	NC-(C)-(C)-CH((CH,)CH-C,H)	PCH-5	NC-(3)-(H)-C1H
83	NC -(3)- N-CH -(3)-C,1H,	РСН-7	NC (
ŷ.	NC-(0)- N *CH-(0)-C+H4	HPE -33	NIC1-(H)-(O)-COO-C1N1
. St X	MC-(O)-(O)-C1HH	нРЕН -23	HIG1-(H)-(Q)-COO -(H)-C1H1
K -21	NC-(C)-(C)-C;H4	нрен-43	H,C,-(H)-(C)-COO -(H)-C,H,
6-5	"c-(C)-(C)-oc'1",	PEPN 4	нс-⊙-∞с-⊙-с•н•
M-15	NC-(()-(0)-00'1H"	PEPN S	NC-(O)-00C-(O)-C1H*
M-21	NC-(2)-(0)-001Hm	PEPN 6	NC-(3)-00C-(3)-C,Hg
M-24	"н°⊃О—О)—ос°н"	PEPN 7	мс-{○}-∞с-{○}-с,нъ
7 - 15	MC-(Q)-(Q)-01Hm	P ₃ 5	NC - (C) - (C) NB
BCH-5	NC-()-()-(+++	P ₃ 7	*#15- \ ___
PCH-3	HC (H-C, H,	T P ₃ 4	MC-Q(C)-(C, M)

† See LC code.

ALIGNMENT OF LIQUID CRYSTALS

(3) Abbreviations used in the text

R-N=N-R'	$R \longrightarrow \bigcap_{i \in I} -N = I_i \longrightarrow I_i$	(pSB)	(XCB)	(XOCB)		$R \leftarrow \bigcirc \bigcirc COO \leftarrow \bigcirc \bigcirc R'$	tiine (nSB) < 0) > 0) xybenzene)	R—H——R'		R CH=N CH=N	$R \longrightarrow \bigcirc \bigcirc C = C \longrightarrow \bigcirc P'$	$R + \left(\bigcirc \right) - C \equiv C - \left(\bigcirc \right) - R'$
LCs with an azo linkage	LCs with azoxy linkage	Azoxys eutectic mixture 4,4'-Bis(n-butoxy)azoxybenzene 4-Butoxybenzylidene-4'-cyanoaniline (pSB) 4-n-Butyl-4'-ethoxy-a-chlorostilbene	Cyanobiphenyls C.H2241	4-Cyanobenzylidene-4'-n-octylaniline $C_4H_{2^{n+1}}$ —O—	Cyano esters Cholesteryl nonanoate 4.4'-Di-n-butylazoxybenzene Dielectric anisotropy	LCs with an exter linkage	4-Methoxy-4'-butylazoxybenzene 4-Methoxybenzylidene-4'-n-butylaniline (nSB) 4-Methoxy-4'-pentyltolan Negative dielectric anisotropy (Δε < 0) Alkoxycyanobiphenyls Positive dielectric anisotropy (Δε > 0) Para-azoxyanisole Para-azoxyphenetol (4,4'-ethoxyazoxybenzene)	Phenylcyclohexane	Mixture of positive LC esters 4-Propyloxy-4'-heptyltolans	LCs with a Schiff base linkage	LCs with a double bond linkage	LCs with a triple bond linkage
Azo	Azoxys (A)	Azoxys eut. mixt. BAB BBCA BECS	CB	СВОА	CE CN DIBAB Le	Esters	MBAB MBBA MPT " OCB PAA PAP	PCH	p Esters mixt. PHT	SB	Stilbenes (S)	Tolans (T)

ing mechanism with reproducibility not guaranteed cooling also favors the LC alignment but such tricks characterize a weak align-

cause of alignment may not be the supposed mechanism. alignment immediately on filling the cell. In the case of weak effects the real For practical purpose a good aligning process has to produce a uniform

A CRITICAL REVIEW OF THE LITERATURE

The alignment of liquid crystals on smooth surfaces

I.1.1 Inorganic Substrates

may help homeotropic orientation, as observed with MBBA, however, acid washed glasses align azoxys parallel to the substrate. ¹⁰ Oxidation of SnO₂ or biphenyls. 25,26 purities on LCs alignment in many compounds. Boiling a glass plate in sulattempts to rationalize the alignment effect observed on 12 different LCs led In₂O₃ coating in an oxygen plasma lead to layers causing parallel alignment of furic acid is supposed to unpolish it, as does dipping in HF solution, 10 which Recent investigations gave elegant evidence of the dramatic influence of impolar molecules tend to induce perpendicular alignment (see paragraph 3). to the hypotheses that it was due to a homeotropic aligning impurity, 'as most tropic alignment on acid treated surfaces is limited to a few Schiff bases, and strate surface is obtained with fired or detergent cleaned glasses. 6.8 Homeotreated glasses⁵ or oxides, ^{6,7} but nonuniform alignment parallel† to the submolecules will align perpendicular to the surface (homeotropically) of acid dures employed in the substrate preparation also play a role, e.g., MBBA alignment is poor as the substrate surface is ill defined. The cleaning proceuid crystals (Table I). The reproducibility and uniformity of this type of The surfaces of glasses, oxides and metals exhibit an aligning influence on liq-

not be obtained from such surfaces sometimes. It has to be considered that reliable homeotropic alignment may tropic alignment using evaporated inorganic layers although it does occur alignment is observed. We have been unable to obtain reproducible homeoserved to induce homeotropic alignment of certain LCs, generally parallel layers have been collected. One sees that, although some layers have been ob-In Table I reported alignment effects of cleaved crystals and evaporated

respectively.

but film uniformity and the substrate used influence the observed results. Polymer coatings on glass substrates can be employed to align liquid crystals,

† The terms "homogeneous and parallel" imply parallel uniform, and nonuniform alignment

I.1.2

Organic polymers

l a	lla	. III P	IV b	V b	VID	VII b	VIII	l b	ПЬ	III a	IV a	V a
LiF ± 12	CaF ₂ × 20 ⊥ 12	***************************************								BN × 19	C × 6,20.24	
NaF × 11 NaCl × 20,90b	MgF ₂ × 15,20,22 Mg(0H) ₂ ⊥ 10 (Brucite)									Al × 17.20 Al.0. × 17.20.21 ± 13.14.20 Aluminosilicate	Si ₁ N ₄ × 18 Si ₀ ₂ × 17, 19,20 es: × 10,23	
KBr × 20,90b KI × 3	(0.000,0)		TiO ₂ × 19 20 22		Cr ₂ 0, × 21 Cr × 17	Fe;0, ± 16	Ni0 × 21		ZnS × 21 ± 13 Zn0 × 11		Ge ₂ O₁ ⊥ 16	AsS, × 10.23
		Y:01 × 19	Zr0 ₂ × 20 ± 20							In ₂ O ₁ × 17.20 ± 17.20	Sn0: × 6,7,20.21 ± 6,7,20	
Cs:0 × 19	BaTiO₄ × 22	CeO ₂ × 19 ThF ₄ ± 13		TaN₁×⊥ 16			P1 × 17,20	Au × 17,20	HgF ₂ × 19		Pb0 ₂ × 21	

Often the same layer has been reported to induce either parallel or perpendicular alignment. Our investigations have generally shown the nematic orientation to be parallel to the surface. Impurities or surface hydrolysis may cause a homeotropic orientation of some LCs which is not characteristic of the surface. ⊥ = homeotropic alignment; × = parallel, nonuniform. Reference 20 to our work means that we have observed these alignments on our test LC mixtures.

TABLE I Attended of name to 1 Co on increasing substrates

ALIGNMENT OF LIQUID CRYSTALS

cated by the case of polytetrafluoroethylene (PTFE) films. As a result of an that some films gave a homeotropic alignment while others did not. An atthat MBBA does not align on PTFE films transferred by sliding contact on to summarizes our results for cast films, together with the available published which is preferably thin in order to avoid an excessive potential drop in the dielectric layer. The film may be transferred to the surface from a liquid, while polymer casting and thermal or plasma polymerization of the monomer have also been used. The most common method is to form the polymer from parexperiment with a series of photopolymerized 1000 Å PTFE films we observed tempt to determine the critical surface tension of these films proved that a continuous variation of the contact angle of liquids of different surface energy served that films obtained from fluorinated polymer suspensions did not give homeotropic alignment of ester or biphenyl mixtures. It has been reported glass substrates,38 while sputtered PTFE films align most LCs, including MBBA. 34 Such a discrepancy is evidently related to the state of the surface of Polyimide layers have been claimed to align MBBA homogeneously²⁷ but our investigation proved that the alignment is generally parallel but not uniform. The alignment properties of most polymers have been evaluated, and Table II data. A wide variety of methods³4 has been used to form the polymer layer tially polymerized solution by dipping or spin coating followed by curing That the condition of film formation determines the final result may be indicould not be obtained on those films which did not align LCs. It was also obthe deposited film.

Smooth, uniform polymer layers of PTFE and silicones induce homeotropic alignment of most commercial LCs with the exception of phenylcyclohexane (Merck Zli 1132) and tolan ³⁴ mixtures. Alignment of these mixtures may be obtained by plasma polymerization of fluorinated alkene monomers ²⁵ or perfluorocycloalkanes ³¹ and by films deposited from silicone solutions (see also paragraph 4). These homeotropically aligning plastic films excepted, other polymers, including plasma deposited films, do not give an uniform parallel alignment and a further rubbing is needed to provide uniformity.

Polymer coatings do not sustain high temperatures and displays employing such coatings must be sealed with an adhesive which is compatible with the polymer layers. This calls for a careful choice of adhesive in the case of silicones. Fluorinated polymers deposited by plasma polymerization of perfluorocycloalkanes, ³¹ as well as polyimide layers ³⁰ are reported to be compatible with a glass frit sealing.

In general, smooth layers of glasses, oxides or polymers orient the nematic director parallel to the substrate but do not induce uniform, reproducible LC alignment.

TABLE II Alignment of nematic LCs on organic polymers

×	MBBA/TN200/ E7/TN103	Natural	Thick film	96-82	Kalle	Suprathen Skyn	Polyalkenes
	COLNIL						b. erp/clene
11	MBBA/TN200/	Вирреф	Thick film	98-36	Kalle	Suprathen Skyn	b. ethýlene
H	MBBA/esters/	Rubbed	Plasma polym.	LT	•••	•••	b. isobutene
11	n. Schiff base mixt.	Knppeq	Polym. sol.	33 (34)	slüH	00E1 lioylo4	p. butadiene
11	n. Schiff base mixt.	Rubbed	Polym. sol.	35-42	movsR insidusiiM	RE 377	Polyactylics p. actylate
II	MBBA/csiers/ . colanes	Кирред	Plasma polym.	33-44		•••	p. methyl methactylate
11		1 11 4	. ,	(4E) TE			
					Taoka Chem.	Ep l	p. cyanoacrylaie
II ×	MBBA/TN103/E7	Rubbed	mli-1	44	JqiW	Вагех	p. actylonitrile
II	MBBA, azoxys,	Кпррса	Polym. sol.	LE	Wackez	Polyviol 13/140	Polyvinyls p. vinyl alcohol
11	n. Schiff base mixt.	Rubbed	Polym. sol.	۲٤	Nichia Paint	novniV	p. vinyl acetate
	LN103, E7					Ut-UZ & IsnivodA	p. vinyl butyral
	TN103, E7					Genotherm	p. vinyl chloride
	TN103, E7	paggnyi	mii-i	66	Kalle	Сепогрегт	p. vinyl chloride
т	2CB: 80CB	Natural	Polym. sol.		•••		p. vinyl pyridinium
11	MBBA/csters/	Rubbed	Poiym. sol.	27-33	Daido Kogyo	Styrite	p. styrenes p. styrene
II.	Schist base mixt	Knppeq	Sol. p. styrene +	•••	•••	•••	
	T		Rubbed MBBA/TN200, Rubbed MBBA/Sesters, Rubbed n. Schiff base mixt. Rubbed MBBA/Sesters, Rubbed MBBA/TN103/E7 Rubbed MBBA, azoxys. Rubbed Azokiff base mixt. Rubbed Schiff base mixt.	Thick film Rubbed MBBA/TN200\ Plasma polym. sol. Rubbed MBBA/csters\ Polym. sol. Rubbed n. Schiff base mixt. Polym. sol. Rubbed n. Schiff base mixt. Polym. sol. Rubbed MBBA/csters\ Polym. sol. Rubbed MBBA, asoxys. Polym. sol. Rubbed MBBA	28–36 Thick film Rubbed MBBA/TN1030, 21. Plasma polym. sol. Rubbed MBBA/TN103, E7 33 (34) 34 (34) 35–44 Plasma polym. sol. Rubbed MBBA/esters/ 35–45 Polym. sol. Rubbed MBBA/esters/ 35–36 Polym. sol. Rubbed MBBA/esters/ 35–36 Polym. sol. Rubbed MBBA, azoxys. 36–37 Polym. sol. Rubbed MBBA, azoxys. 36–38 Polym. sol. Rubbed MBBA, azoxys. 36–39 Polym. sol. Rubbed MBBA, azoxys. 37 Polym. sol. Rubbed MBBA, azoxys. 38–34 Polym. sol. Rubbed MBBA, azoxys. 39–34 Polym. sol. Rubbed MBBA, azoxys.	Kalle 28–36 Thick film Rubbed MBBA/TN1004	Supraithen Skyn Kalle 28–36 Thick film Rubbed MBBA/esters/

Polymer	Commercial name	Source	10 ⁻³ J m ⁻²	Mode of coating	Surface state	Liquid crystals	Alignment	Ref.
Polyparaxylylene								
p.(p. xylylene)	Parylene	(Union Carbide) (Backelite)	•••	Vapor polym.	Rubbed	DIBAB	11	29
D. I		(BXC)	•••	Vapor polym.	Rubbed	5CB	11	34
Polyesters, polyurethanes p. esters	1	.						
p. esters	Lumi rror Mylar A, B, C	Toray Dupont	40-43 40-43	Polym. sol. Thick film	Rubbed Natural	Schiff base mixt. MBBA/TN200/ TN103, E7	×	20
p. esters	Mylar A, B, C	Dupont	40-43	Thick film	Rubbed	azoxy ent. MBBA/TN200/ TN103, E7	11	20
p. esters	Melinex	ICI	40-43	Thick film	Natural	azoxy ent. MBBA/TN200/ TN103, E7	×	20
p. esters			40-43	Thick film	Rubbed	azoxy ent. MBBA/TN200/ TN103, E7	11	20
p. esters	Hostaphan	Kalle	40-43	Thick film	Natural	azoxy ent. MBBA/TN200/ TN103, E7	×	20
p. esters	Hostaphan	Kalle	40-43	Thick film	Rubbed	azoxy ent. MBBA/TN200/ TN103, E7	11	20
p. urethane	V. Chroma	Daï Nippon Toryo		Polym. sol.	Rubbed	azoxy ent. Schiff base mixt.	11	36
Polyamides p. amide	Capran	All Chem.	33-42	Thick film	Natural	TN200, TN103, E7	×	20
p. amide	Versamid	0 14"			Rubbed	TN200, TN103, E7	11	20
p. methylamide (Nylon 6)	Sumitherm	Gen. Mills Sumi Denks	33-42	Sol. in LC Polym. sol.	Natural Natural	MBBA Schiff bases	×	56 36
olyimides p. imide	Nolimid	Rhône Poulenc		From monomer sol.	Rubbed	TN200, TN103, TN104, E7	II	20
•								
p. imide	Nolimid	Rhône Poulenc	•	From monomer sol.	Rubbed	E8, ZLi1132	11	20
p. imide p. imide	Nolimid Nolimid	Rhône Poulenc Rhône Poulenc	•••	From monomer sol. From monomer sol.	? Rubbed	MBBA Schiff bases, CB,	11	27
p. imide	Kapton	Dupont		Thick film	Natural	Esters azoxys, azo TN200, TN103,	∥ ×	30,31 20
p. imide	Kapton	Dupont		Thick film	Rubbed	TN404, E7, E8 TN200, TN103,	×	20 20
•	Kapton	Бироп		11100 11111	Nuovia .	TN404, E7, E8		
olysulfone p. sulfone	S 179	Appl. Sc. Lab	41	Polym. sol.	Natural Rubbed	TN103, E7, azoxy n. esters mixture	×	20 20
olyheterocycles p. quinoxaline	. •••	Monomer		Polym. sol.	Rubbed	p. nematic mixt.	11	28
p. benzthiazole		Monomer	•••	Polym. sol.	Rubbed	p. nematic mixt.	II	28
p. benzoxazole	•••	Monomer	•••	Polym. sol.	Rubbed	p. nematic mixt.	II	28
ellulose derivatives Methyl cellulose	•••	Shinetsu Kagaku	35-42	Polym. sol.	Rubbed	Schiff base mixt.	1	36
65SH4000 Cellulose acetate	•••	Ceta SA	39	Thick film	Natural	MBBA/TN103/	×	20
Cellulose acetate		Ceta SA	39	Thick film	Rubbed	E7/azoxys MBBA/TN103/	II	20
Cellulose acetate Cellulose triacetate	LT105 LT105	Daicel Daicel	39 39	Polym. sol. Polym. sol.	Rubbed Rubbed	E7/azoxys Schiff base mixt. Schiff base mixt.		36 36
and various esters Nitrocellulose Nitrocellulose	HIG40 Celluloid	Asai Kasei	34-42 34-42	Polym. sol. Thick film	Rubbed Natural	Schiff base mixt. TN200/TN103/ E7/azoxys	 X	36 20
Nitrocellulose	Celluloid	•••	34-42	Thick film	Rubbed	TN200/TN103/ E7/azoxys	11	20
olysilanes p.(glycidopropyl-		•••	41 (34)	Dipping in monomes	Nat ?	MBBA, esters,	11	34
trimethoxysilane) p. hexamethylsiloxane	•••		24 (34)	sol. Plasma polym.	Natural	tolans, CB MBBA, esters,	Τ	34
Silicone rubber	KE45	Shinetsa Kagaku	22-24	Polym. sol.	Natural	tolans, CB Schiff base mixt.	T	36
Silicone rubber	CAF RTV	Rhône Poulenc	22-24	Polym. sol.	Natural	TN200/103-E7, azoxys	Ī	20

Ref.	Alignment	Liquid crystals	Surface	Mode of coating	10-1 J m-2	Source	Commercial name	Polymer
50	Т	TN200/103-E7,	Natural	Polym. sol.	77-74	Chemie Moudon	Elcon 111	Silicone tubbet
70	Т	TN200/103-E7,	Natural	Thick film	77-24	Frabera	•••	p. ester silicone
16	Т	WBBY\EBBY szoxks	Natural	Dipping	55-54	•••	•••	p. methylphenylsiloxane
16	Ţ	MBBA/EBBA	Natural	amqqrd BniqqiQ	22-24	•••	•••	p. methyl p. dimethylsiloxane
								.ixim
98	'	p. S. Bműt	Natural	Polym. sol.	77-74	onelwod enoda	900 !5	p. methylsiloxane
70	т	<i>L</i> 3	Natural	·los		Rhone Poulenc	966 !S	p. methyl, phenyl (45%) siloxane
70	×	103	Natural	JoS.		Rhône Poulenc	966 !S	p. methyl, phenyl (45%) siloxane
70	×	7611	IstutaN	Sol.	•••	Rhône Poulenc	966 !S	p. methyl, phenyl (45%) siloxane
34	Т	MBBA, esters.	Natural	Sputtering	16-22	JuoquQ	Teflon	Fluorinated polymers p. tetrafluoroethylene
229	11	CB colans	Natural	Plasma polym.	75 (34)	•••	•••	p. tetrashuoroethylene
8£	×	MBBA, Chlorostilbene,	Natural	Sliding contact	(8£) 91			p. tetrashuoroethylene
50	×	azoxys CB, TN103	Natural	Photo polym.	not meas-	•••	***	p. tetrafluoroethylene
50	×			Thick film	urable 16.51	Dupont		p. Auoropropyl Co
υc	11		poqqua	Thick film	32.3	taoanG	Tedlar	tetrafluoroethylene FEP p. vinylidene fluoride
35 30	T II	Schiff base mixt.	Rubbed	Plasma polym.	(35) 81	Dupont		p. 1,3-dimethyleyelohexane
35	Ţ	Schiff base mixt.	Natural	Plasma polym.	55-54 (35)	•••	• • •	p. perfluorocyclohexene
35	T	Schiff base mixt.	Natural	mylog smasslq	(35) 22-02	•••	•••	p. perfluoro
LE	II-T	Schist base mixe.	Natural	16-81	•••	•••	•••	dimethylcyclobutane p. acetylene Co perfluoro-1,3- dimethylcyclohexane
98	11	Schiff base mixt.	Rubbed	Polym. sol.	19	uskoT idsidustiM	•••	stanton (spinos)
	n							bioiv

^a Critical surface tension from Polymer Handbook. 2nd ed., edited by J. Brandrup and E. H. Immergut. (Wiley, New York, 1975), except as otherwise indicated.

⊥ = homeotropic alignment (perpendicular to glass surface); || = alignment, uniform, parallel to glass surface; • = molecular alignment too difficult to distinguish between perpendicular and parallel alignments and to obtain consistent experimental results; ||√L = molecular alignment changed near the phase transition temperature from the parallel alignment at lower temperatures to the perpendicular alignment at higher temperatures:

X = parallel nonuniform.

ALIGNMENT OF LIQUID CRYSTALS

I.2 Alignment on grooved surfaces

Rubbing, tangential evaporation or shallow angle ion beam etching (SAIBE) produce a wavy surface. It has long been acknowledged that the rubbing of glass plate induces uniform alignment of LCs, ^{39,40} with the nematic director nearly parallel to the substrate surface. ⁴¹ For some time it was considered that the material used for rubbing determined the efficiency of the process.† It is now clear, however, that, as proposed many years ago, ³⁹ any rubbing material, e.g., paper, tissues, brushes and polishing powders, gives good results. Some LCs align more easily than others and reproducibility is not very good on substrates that are simply rubbed. The use of polishing powder, such as diamond paste, improves the alignment and is a requisite for hard layers of silica or fluoride.

Glass has a surface layer extending one micron in depth which has higher entropy than the bulk⁴² and may be easily deformed by rubbing, producing a wavy surface. Because of its higher energy this surface layer melts 125°C below the softening point of the glass⁴² (i.e., around 475°C for soda lime glass) and, therefore, the effects of rubbing disappear above this temperature. Glass frit sealed cells, therefore, require the use of layers of high melting point compounds. SiO₂ deposited by sputtering, CVD or an electron gun, as well as MgF₂. ^{15,20} BaTiO₄. TiO₂. ²² Cs₂O, Y₂O₃, BN, ¹⁹ CeO¹⁹ or Si₃N₄. ¹⁸ layers give, after rubbing with diamond paste or cerium oxide powder, ^{43,20} good, homogeneous aligning layers but the results are very sensitive to further processing. It is likely that these hard materials are difficult to groove sufficiently.

Oxides, ¹⁷ fluorides and metal layers evaporated obliquely to a substrate generally align the LC molecules parallel to, or at a slight angle from, the surface ⁴⁴ but tangentially evaporated calcium fluoride is reported to align the molecules nearly perpendicular ¹² although we could not reproduce this result. Slant evaporation of a metal and subsequent thermal oxidation ²¹ or shallow angle ion beam etching (SAIBE) ^{45a} produce the same results. Obliquely evaporated SiO_x and MgF₂ layers give reproducible results and this method is widely used for the fabrication of small displays; it is further compatible with any type of sealing.

When high temperature resistance is not required, as in experimental cells or plastic sealed displays, it is advantageous to coat the electrodes with a soft polymeric layer which is rubbed afterwards. Any polymer is suitable, pro-

alignment depends on the polymer layer thickness and the substrate unialign homogeneously. Rubbed polyvinyl alcohol or polyvinyl butyraldehyde vided that it is possible to deposit it as a thin film (Table II) and most LCs will or of SiO₂ 51 induces homogeneous alignement of a LC. 47,50 Many surfaces a piece of polymer, have also been described as producing good aligning rubbing of substrates on which a polymer powder was spread, or rubbing with wide variety of methods has been used to form films on a glass substrate. The diate layer that promotes adhesion and uniformity. As previously described, a formity. It is often advantageous to provide the electrodes with an intermeing layer simultaneously, 33 but the polarizing effect is weak. The quality of the have been proposed as a method of forming an internal polarizer and an alignwhich has led to the claim that any striated surface, whatever the substrate and the method used to produce them, 55 induces homogeneous alignment. In fact been shown by photomicrographs. A grating made of a photoresist layer 47,34 paste^{43,47} and the striated nature of tangentially evaporated layers⁴⁸⁻⁵³ has been clearly shown, the grooves produced by rubbing with a polishing layers. 46 Although the surface deformation of glasses and polymers has not tend to align LC's parallel and striations on these surfaces provide uniformity, polymethylsiloxane still promotes an homeotropic alignment after it has been homeotropic aligning layers, e.g., evaporated CaF_2 , show at most a leaning of rubbed (see Figure 9). the molecular axis when striations are induced by oblique evaporation, and

One point which is not clear is the surface nature of evaporated inorganic layers which are sometimes found to induce homeotropic alignment (not in a reproducible manner) but, once rubbed, promote a homogeneous alignment.

1.3 Alignment by surface active agents

It has long been observed⁵⁶ that surface active agents promote LC alignment. They may be either dissolved in a LC, or deposited on the cell walls. Small amount of surfactants may be dissolved in LCs conveniently through a common solvent which is evaporated afterwards. Spontaneous homeotropic alignment of the mixtures is observed on glass or oxide surfaces. Most surfactants have been described as effective (Table III) and examples are given for negative LC mixtures. These observations have been hastily extended to "liquid crystals" in general. As surfactant induced alignment is complex depending on the substrate, mode of application, LC composition (see § II.1.8), the use of these data needs critical evaluation. ^{57,65} For displays operating in the field effect mode, the increase in conductivity due to an ionic dopant is a disadvantage and, therefore, nonionic compounds are preferred. Cationic surfactants, which are long chain substituted ammonium salts, are very effective in promoting homeotropic alignment of negative ⁵⁸⁻⁶⁰ and positive ^{20,117} LCs although they increase the LC conductivity. Figures obtained for various con-

[†] Alignment of LCs by rubbed surfaces is sometimes wrongly attributed to J. F. Dreyer who omitted, in his paper "Orientation of the surface of glass" [Glass. Ind. 29, 197 (1948)], to quote the work of Zocher and Coper (Ref. 39). This point has been cleared up in a letter to the editor by C. D. West [Glass. Ind. 30, 272 (1949)], which provides an English translation of the main paragraph of Ref. 39. P. Chatelain, ⁴⁰ who studied in detail LC alignment by rubbed surfaces, also ignored this work which clearly established the influence of rubbed glass surfaces on the orientation of PAA (which was used to prove the glass surface anisotropy) and that it was a property of the clean surface.

TABLE III

Surfactants agents reported to induce homeotropic alignment of nematic LCs

Surfactants	Liquid crystal	Ref.
I. ANIONIC		
$R = C_n H_{2n+1}$ $n > 10$ $n = 8, 11, 19$	MBBA n-Schilf base mixt.	\$ 61
$R = C_1H_{12}CH=CH-(CH_2), \text{ (oleic acid)}$ $R = C_nF_{2n+1}$ $R = C_nF_{2n+1}$ $R = C_nF_{2n+1}$ $R = C_nF_{2n+1}$	n-Ester mixt. MBBA MBBA Esters	3 00 00
$R = C_n H_{2n+1}$ $n = 18 \text{ (octadecylmalonic acid)}$		
1.2. AROMATIC ACIDS: $R \leftarrow \bigcirc$ —COOH	n-Schiff base mixt. n-Ester mixt.	
$R = 4NH_{3}; 4 \text{ OH}; 3 \text{ OH}$ $R = C_{n}H_{2n+1}O$ $n = 9, 10, 12, 14, 16, 18$ $R = C_{n}H_{2n+1}$ $R = C_{n}H_{2n+1}$ $R = C_{n}H_{2n+1} - COO$ $R = 3, 17$	n-Azoxy mixt.	61 62 66 62
$R \longrightarrow CH_1COOH$ $R = 4NH_1, 4 OH, 2-5 OH$	n-Schiff base mixt. n-Ester mixt. n-Azoxy mixt.	
$R \longrightarrow (CH_1)_2 \longrightarrow COOH$		
R = 4NH ₃ , 4 OH, 3-4 OH	n-Schiff base mixt. n-Ester mixt. n-Azoxy mixt.	19
В───СН=СН—СООН	n-Azoxy mixt.	62
$R = C_n H_{2n+1}O$ $n = 10, 12, 18$ $R = C_n H_{2n+1} - COO$ $n = 1, 12, 18$	n-Azoxy mixt.	
. В Сооон		
$\begin{aligned} R_1 &= C_0 H_{11}O & R_2 &= CI \\ R_1 &= C_1 u H_{13}O & R_2 &= CI \\ 1.3. & \textit{CARBOXYLIC ACID WITH LC STRUCTURE.} \end{aligned}$	n-Ester mixt.	99

1.3.1. Derivatives of Schiff's base

ALIGNMENT OF LIQUID CRYSTALS

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TABLE III (continued)

Surfactants	Liquid crystal	Ref.
$R' = H$, $R_n = 4$ OH, 3 OH, 2 NO ₂ -3 OH, 3-4 OH, 2-4 OH, 2-4-6 OH $R_n = C_n H_{2n-1}O$ $n = 1, 7, 13$, $R_n = C_n H_{2n-1}O$ $n = 1, 7, 13$ $R' = OH R_n = C_n H_{2n-1}O$ $n = 0, 1, 3, 7, 13$	n-Schiff base eut. mixt. n-Esters eut. mixt. n-Azoxy eut. mixt. n-Azo eut. mixt.	61, 62
$R_{n} = \bigoplus_{\substack{A = 4 \text{ OH. } 3 \text{ OH. } 3-4 \text{ OH} \\ R_{n} = C_{n}H_{2n} \text{ IO}}} - CH = N \bigoplus_{\substack{A = 1, 3, 4, 15 \\ R_{n} = C_{n}H_{2n}}} - CH = N \bigoplus_{\substack{A = 1, 4, 7, 13 \\ R_{n} = C_{n}H_{2n}}} - CH = N \bigoplus_{\substack{A = 1, 4, 7, 13 \\ R_{n} = C_{n}H_{2n}}} - CH = N \bigoplus_{\substack{A = 1, 4, 7, 13 \\ R_{n} = C_{n}H_{2n}}} - CH = N \bigoplus_{\substack{A = 1, 4, 7, 13 \\ R_{n} = C_{n}H_{2n}}} - CH = N \bigoplus_{\substack{A = 1, 4, 7, 13 \\ R_{n} = C_{n}H_{2n}}} - CH = N \bigoplus_{\substack{A = 1, 4, 7, 13 \\ R_{n} = C_{n}H_{2n}}} - CH = N \bigoplus_{\substack{A = 1, 4, 7, 13 \\ R_{n} = C_{n}H_{2n}}} - CH = N \bigoplus_{\substack{A = 1, 4, 7, 13 \\ R_{n} = C_{n}H_{2n}}} - CH = N \bigoplus_{\substack{A = 1, 4, 7, 13 \\ R_{n} = C_{n}H_{2n}}} - CH = N \bigoplus_{\substack{A = 1, 4, 7, 13 \\ R_{n} = C_{n}H_{2n}}} - CH = N \bigoplus_{\substack{A = 1, 4, 7, 13 \\ R_{n} = C_{n}H_{2n}}} - CH = N \bigoplus_{\substack{A = 1, 4, 7, 13 \\ R_{n} = C_{n}H_{2n}}} - CH = N \bigoplus_{\substack{A = 1, 4, 7, 13 \\ R_{n} = C_{n}H_{2n}}} - CH = N \bigoplus_{\substack{A = 1, 4, 7, 13 \\ R_{n} = C_{n}H_{2n}}} - CH = N \bigoplus_{\substack{A = 1, 4, 7, 13 \\ R_{n} = C_{n}H_{2n}}} - CH = N \bigoplus_{\substack{A = 1, 4, 7, 13 \\ R_{n} = C_{n}H_{2n}}} - CH = N \bigoplus_{\substack{A = 1, 4, 7, 13 \\ R_{n} = C_{n}H_{2n}}} - CH = N \bigoplus_{\substack{A = 1, 4, 7, 13 \\ R_{n} = C_{n}H_{2n}}} - CH = N \bigoplus_{\substack{A = 1, 4, 7, 13 \\ R_{n} = C_{n}H_{2n}}} - CH = N \bigoplus_{\substack{A = 1, 4, 7, 13 \\ R_{n} = C_{n}H_{2n}}} - CH = N \bigoplus_{\substack{A = 1, 4, 7, 13 \\ R_{n} = C_{n}H_{2n}}} - CH = N \bigoplus_{\substack{A = 1, 4, 7, 13 \\ R_{n} = C_{n}H_{2n}}} - CH = N \bigoplus_{\substack{A = 1, 4, 7, 13 \\ R_{n} = C_{n}H_{2n}}} - CH = N \bigoplus_{\substack{A = 1, 4, 7, 13 \\ R_{n} = C_{n}H_{2n}}} - CH = N \bigoplus_{\substack{A = 1, 4, 7, 13 \\ R_{n} = C_{n}H_{2n}}} - CH = N \bigoplus_{\substack{A = 1, 4, 7, 13 \\ R_{n} = C_{n}H_{2n}}} - CH = N \bigoplus_{\substack{A = 1, 4, 7, 13 \\ R_{n} = C_{n}H_{2n}}} - CH = N \bigoplus_{\substack{A = 1, 4, 7, 13 \\ R_{n} = C_{n}H_{2n}}} - CH = N \bigoplus_{\substack{A = 1, 4, 7, 13 \\ R_{n} = C_{n}H_{2n}}} - CH = N \bigoplus_{\substack{A = 1, 4, 7, 13 \\ R_{n} = C_{n}H_{2n}}} - CH = N \bigoplus_{\substack{A = 1, 4, 7, 13 \\ R_{n} = C_{n}H_{2n}}} - CH = N \bigoplus_{\substack{A = 1, 4, 7, 13 \\ R_{n} = C_{n}H_{2n}}} - CH = N \bigoplus_{\substack{A = 1, 4, 7, 13 \\ R_{n} = C_{n}H_{2n}}} - CH = N \bigoplus_{\substack{A = 1, 4, 7, 13 \\ R_{n} = C_{n}H_{2n}}} - CH = N \bigoplus_{\substack{A = 1, 4, 7, 13 \\ R_{n} = C_{n}H_{2n}}} - CH = N \bigoplus_{A = 1$	As above	29
$R = C_{c}H_{3r-1} $ $R = C_{$		
$R = C_k H_{17}O$	n-Ester mixt.	26
$R \longrightarrow CH = N \longrightarrow CH = C \longrightarrow CH = C \longrightarrow COOH$ CH_3 $R = CH_3O \qquad R = C_3H_3O \text{ (slow)}$ 1.3.2. Derivatives of LC azoxy structure	n-Ester mixt.	. 95
$R = C_n H_{3n_1}O$ $R = C_n H_{3n_2}O$ $R = C_n H_{3n_3}O$ $R = C_n H_{3n_3}O$ $R = 15$ $R_n = C_n H_{3n_3}$ $R = 14$	n-Azoxy cut, mixt.	62
$R = C_n H_{3r_1}O$ $R = C_n H_{3r_1}O$ $R = 2.4$ 1.3.3. Derivatives of LCs with exter structure	n-Azoxy eut. mixt.	62
в—Соо—Соон	n-Azoxy mixt. n-Ester cut. mixt.	62

Surfactants

 $R = C_n H_{2n+1}O \qquad n = 10, 14$ $R = C_n H_{2n+1} \qquad n = 4, 8, 18$ $R = C_n H_{2n+1}COO \qquad n = 3, 15, 17$

1.3.4. Derivatives of LCs stilben structure

—CH=CH-СООН

1.3.5. Derivatives of LCs azo structure

 $R = C_n H_{2n+1}O$ n = 8, 18 $R = C_n H_{2n+1}COO$ n = 6, 9

−сн=сн-(() }-соон

TABLE III (continued)

ALIGNMENT OF LIQUID CRYSTALS

n-Schiff base mixt.		n-Schill Odac HIIAL.	n-Schiff have mixt	PAA MBBA-E7, TN103, ZLi1132	MBBA	MDBA			MBBA MBBA	MBBA	E7		a-AZO EUL MIXL			Chloro stilben mixt.	n-Azoxy mixt.	n-Azoxy eut. mixt.			Liquid crystal
70	62	70 39	70 60	58 58 75-20	58.57	;			58 58	\$8	45c		62	;			62	62			Ref.
$C_nH_{2n+1}-NH_2$ n > 12 n = 16 n = 18	$C_n H_{2n+1} - HN - CO - NH_2$ $n = 16$	$C_n H_{2n+1} - CN$ $n = 7$	ת = 11 R = 6	$R - COO - C_n H_{2n-1}$ $n = 8.18$ $R = CH_3$	3.1. ALIPHATICS ESTERS, NITRILES, UREA, AMINES OR ALCOHOL	3. NONIONIC	×2/ 0/	R-C XiOH	2.5. CARBOXYLATOCHROMIUM COMPLEXES	$R' = NH_2$, NO_3 , Cl , H , CH_3 , C_3H_3 , C_4H_5 , C_6H_{13}	$C_{16}H_{11}-N^{2}$ 000C $-\left(\bigcirc\right)-R^{2}$	2.4. 7-ALKYL-I.8-DIAZABICYCLO[5.4.0]UNDECENAMMONIUM	$n = 12 \qquad \bigvee_{N' - C_n H_{2m+1} X}$		2.3. ALKYLISOQUINOLINIUM SALTS	n = 16, 12 $(X' = Br')n = 16 (X' = (C_4H_3)_4B')$	N^{-} C _n H _{2n-1} X	2.2. ALKYLPYRIDINIUM SALT	(C ₁₁ H ₁₇) ₂ N'(CH ₃) ₂ -(C ₆ H ₅) ₄ B'	2.1.3. X' = tetraphenyl borate	Surfactants
MBBA MBBA MBBA	MBBA	MBBA	n-Schiff base mixt.	n-Schiff base mixt.	<i>чтсонот</i>			p-Schiff base	n-Schiff base mixt. CB	n-Azoxy mixt. n-Ester mixt.	n-Schiff base mixt.	NUM	n-Schiff base mixt.	PAA		n-Ester mixt.	n-Schiff base mixt. n-Ester mixt.		n-Schiff base mixt.		Liquid crystal
5 75 60	66	60	61	74					73			70	54	54	;	68	59 59		69		Ref.

2.1.1. $X^* = halogen$

2.1. ALKYL AMMONIUM SALTS C,H;,-I-N'(R) X

2.1.2. X' = benzoate

C'''H'''N. C'H'

)—0C,H,,

C4H4-N,-(CH1),000C-() -C4H14

n > 10
n = 16 (Cetyl trimethyl ammonium bromide)
C₁F₁SO₂NH(CH₂)₁N'(CH₃)₁

Cobalt, zinc naphthenate Sulfated alcohols Sulfated ethers

2. CATIONIC

1.4. ANIONIC SURFACE ACTIVE AGENTS

1.3.6. Derivatives of cyanobiphenyls

 $R = C_n H_{2n+1}O \qquad n = 1, 6, 12, 18$ $R = C_n H_{2n+1} \qquad n = 6, 10, 14$ $R = C_n H_{2n+1}COO \qquad n = 3, 7, 15$

 $\left\langle \left(\right) \right\rangle$ N==N $\left\langle \left(\right) \right\rangle$ COOH

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TABLE III (continued)

Surfacianis	Liquid crystal	Ref.
C_F_MCF=CF(CH1)1NH(CH1)1N(CH1)1	MBBA/EBBA	92
n = 0 $C_nH_{2n-1} - OH \text{ (amine catal.)}$ $n = 12.14 \cdot 16.18$ $C_nF_{2n-1}(CH_2)_nOH$ n = 7. m = 3	n-Ester eut. mixt. MBBA MBBA/EBBA	76 66 92
3.2. AROMATIC ACID ESTERS		
H ₂ N-(CH ₂) _m -COOC _n H _{2m1}	n-Schiff base mixt.	63
n = 6 to 16 $m = 0, 1, 2$		
$X_a \longrightarrow COO \longrightarrow C_aH_{2m-1}$	n-Schiff base mixt.	63
$n = 4 \text{ to } 12$ $X_n = 4 \text{ OH}$ $n = 2 \text{ to } 18$ $X_n = 1, 2, 6 \text{ OH}$		
3.3. PHENOLS, AROMATIC AMINES		
2.6 Di-r-butyl-4-methylphenol Hydroquinone 4butylpyrocatechin 2-r-butylhydroxyanisole Nisopropyl-A-phenyl p. penylenediamine Phenyl-8-naphthylamine p-hydroxyphenyl-8-naphthylamine	MBBA MBBA MBBA MBBA MBBA MBBA MBBA	122222
3.4. NONIONIC SURFACTANTS		
Epoxy resins (tow molecular weight) Polyamid resins (tow molecular weight) Alkyl phenyl ethers Polyoxyethylated glycols Fluoro polymers	MBBA MBBA MBBA MBBA MBBA	58 50 67 67
4. AMPHOLYTICS		
Lecithin egg Lecithin egg Lecithin	MBBA E7. E8. TN103. TN404. 1132	79-6 20
a Lecithin CH_2 — CH — CH_2O - PO_3 $\bigcirc (CH_3)_2N'(CH_3)_3$ R_1 R_2	PCH, Schiff bases, azoxy, esters	\$9

centrations of cetyl ammonium bromide in a positive ester mixture ROTN 103 are given in Table IV.

The use of these agents has, therefore, been proposed for dynamic scattering mode displays. 46.59 Alignment of positive LCs is also observed with these compounds. Alkylvinyl pyridinium halides provide stronger orientation than ammonium derivatives, being able to align even smeetic phases. In fact most

ALIGNMENT OF LIQUID CRYSTALS

TABLE IV

Alignment and conductivity (o) of p. ester mixture TN103, with added CTAB

	10-8	11
,	7.4×10^{-8}	7
0.5	2.9×10^{-9}	-1
0.25	1.7×10^{-9}	•
0.1	ှိ ဗ	<u></u>
0.01	2×10^{-10}	=
0	1.8×10^{-10}	===
% CTAB	ь	LC alignment

LC. Fatty nitriles, amines, 60 acids 6,61 and esters 61 align MBBA. The effect of nitriles and esters is weak, depends on the substrate, and fails to align biphenyl and ester mixtures. Substituted benzoic acid^{61,62}, its ester, phenols 15,52,54 and octadecyl malonic acid64 are also effective, but branched isopalmitic acid does not align a LC. 65 Derivatives of benzoic acids which possess a liquid crystal structure have been thoroughly investigated 61.62 and a critical study 57 has assessed the more effective dopants. Such compounds, being frequently present as impurities or decomposition products in a LC, cause wrong assignment of he influence of aligning agents or prevent accurate correlations between LC structures and surface treatments. Their effect appears stronger when they are ormed in the cell by LC degradation than when added to the LC. Effective concentrations of additives to the LC are in the range 0.5% to 2%. At smaller concentrations, the alignment presents defects, and higher concentrations ower the nematic to isotropic transition temperature significantly. The main drawback of this method of alignment is that on filling a cell provided with only one fill port, as preferred in industry, the additives absorb strongly in the neighborhood of the aperture and the liquid crystal at the end of the cell contains a lower concentration of additive, producing defects and a conductivity gradient across the cell. Attempts have been made to treat the cell walls before filling. Most of the surfactants described above for doping LC mixtures are effective, as the dissolved surfactant aligns the LC through absorption on the compounds of this type, dissolved in LCs, favor homeotropic alignment of the cell walls.

The orientation of the LC depends on its molecular structure: while most LCs are homeotropically aligned by adsorbed lecithin, azoxy derivatives and 4-methoxypropylcarboxybenzylidene aniline are not aligned. 65 The packing density of the adsorbed amphiphilic layer also plays a role. While homeotropic alignment of MBBA on glasses covered with hexadecyl(=cetyl)trimethyl ammonium bromide (CTAB) occurs at high coverage, 8 lecithin's orienting power decreases with increasing packing density. 65 Fatty amines and alkyl substituted ammonium derivative 66 chains with a carbon content over 10 promote homeotropic alignment, while shorter chains orient the nematic director at an angle from the surface, smaller angles being observed for shorter chains. Heating destroys the aligning layer and cells provided with one fill port must be treated with a solution of the surfactant in a volatile solvant, rinsed,

dried and filled. Although often used in experimental work with lecithin in ether, or CTAB in alcohol, this method has not found industrial acceptance.

These methods give good results on many substrates except plastic coatings but long term stability is not obtained as the absorbed layer slowly dissolves in the LC.

boxylatochromium complexes⁷³ or silanes which are substituted with one long chain polar group, ⁷⁷ e.g., N-octadecyl-N,N-dimethyl-3-aminopropyltri-A large increase in LC conductivity, however, is observed when these commethoxysilane aligns E7 and ROTN 103 homeotropically. Chromium com-MBBA and N-p-(cyanobenzylidene)-p-N-octylaniline homogeneously after rubbing, 77 but we found that the very similar N-ethyl-3-aminopropyltrimethoxysilyl chloride. We have checked that the aligning power is strong for chloride⁷² dissolved in the LC produces homeotropic alignment, probably due covered electrodes catalyzed by amines^{45b} seems to depend on the quality of sorbed on cell walls from a solution has been claimed⁶⁹ to improve the alignthe surface. Baking octadecyltrimethyl ammonium tetraphenyl boride abcloalkanes (crown ethers) improve homogeneous alignment on most oxides.80 homeotropic alignment is observed for positive mixtures. 78 Diazapolyoxycyplexes of dicarboxylic acids align negative LCs homogeneously, whereas acids added to LC mixtures are claimed to impart horizontal orientation.76 mixtures. Few additives, however, promote planar alignment. Dicarboxylic compounds generally induce homeotropic alignment of most practical LC plasticizers: long term stability has not yet been evaluated. Long chain polar by selective treatment of the walls. Nevertheless, these compounds, being way of realizing both homogeneous and homeotropic domains in the same cell most positive nematic mixtures and effective on rubbed layers, providing a to reactions with the electrode surface. The best results are obtained with carited by CVD or solutions of organosilicon compounds. Tributyltin oxide" or the silica layers as we did not obtain reproducible results on silica layers deposment of the negative Schiff's bases. Reaction of fatty alcohols with silica pounds are used. Also, surfaces treated with N-methyl-3-aminopropyltrimethoxysilane align ionic, increase slightly the LC conductivity even with thoroughly cleaned surfaces. Surfactants may also be included in a polymer layer, where they act as Attempts have been made to bind the surface active molecule covalently to

It is interesting to note that reacting tin oxide layers with long chain alcohols ($C_nH_{2n+1}OH$, n=1 to 20), triethanolamine, phenols, cyclohexanol, phenyl acetic acid or valeric acid under pressure promotes homogeneous alignment of LCs as opposed to previous observations. An example is given for tin oxide treated either with n-decanol or cinnamic alcohol on an alkyloxyphenylpyrimidine mixture.

I.4 Alignment by silane treated surfaces

Alkoxys and chlorosilanes interact strongly with glass surfaces. The proposed mechanisms are either reaction with surface silanol groups, or hydrolysis of the silane to a silanol which will further condense into a linear polysiloxane layer (Figure 1). Thus silanes are sometimes considered as surface active agents and sometimes as polymer formjing compounds. As they are of interest for LC alignment they justify a special paragraph.

substrate is exposed to silane vapor, for low boiling point compounds, or silanes. 72a In the case of quaternary ammonium sylil compounds these acetic in water, DMF or acetone. Water solutions are stable for only a few observed that depositing a drop of pure silane on a spinning substrate often tions⁸³ improved the efficiency of the treatment. Silanation also occurs if the methods gave poor results with most LCs and dipping in hot (75°C) soluhours (except aminosilanes). After rinsing with the solvent used to dissolve the hr., (generally 5 min) in 1%-5% solution of the silane in water, toluene, dilute gave better results than dipping in silane solutions. Hydrolysis of the surface treated in a refluxed solution of silane in toluene, for others. We have often tropically can only be persuaded to align a wide range of LCs uniformly and and biphenyls homeotropically.87 Those silanes which do not align homeochlorosilane. Stearyltrichlorosilane has been reported to align Schiff's bases ilar ethyl substituted derivatives, and methyltrichlorosilane or trimethylbeen tried, monodimentyldichlorosilane shows a strong effect, unlike the simdeposition and the nature of the LC. Among the different silanes which have results. It can be seen that the effect of the silane layer depends on the mode of biphenyl mixture E7, is presented in Table V together with some published efficiency for the alignment of the positive ester mixture ROTN 103, and the before coating is sometimes recommended. An evaluation of silane alignment merization of silanes shows that only methylpolysiloxane (and for some LCs, oxane surface layer. Deposition of silicones which are formed by bulk polymost silanes influence the LC alignment by the in situ formation of a polysilparallel to the substrate, after rubbing, as has already been noted. It seems that to form an initial surface layer by decomposition of an organometallic comcompounds. A substrate may alter the film properties and it is advantageous promote a planar alignment, contrary to certain reported observations.86 In methylphenylsiloxane) aligns LCs homeotropically, while the remainder pound before silane treatment. Silylated surfaces degrade upon heating and pounds. A substrate may alter the film properties and it is advantageous to bind a long chain to the substrate, as with stearylsilane or dodecylammonium fact, silanes may be used in two ways: either to form a polysiloxane layer or to form an initial surface layer by decomposition of an organometallic com-Surface treatments with silanes have been effected by dipping during 5 s to 1 25

ALIGNMENT OF LIQUID CRYSTALS

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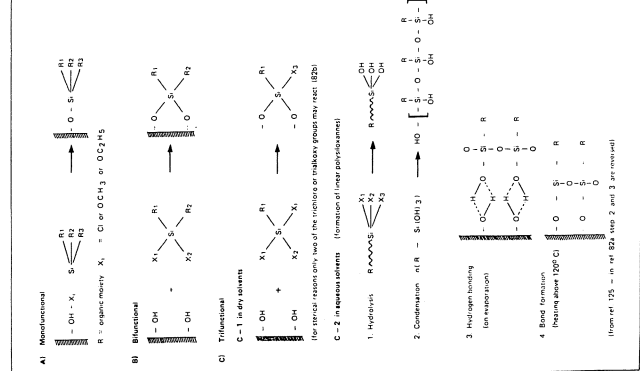
103 gnirus D-08 5 min. Sor in benzene

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apor phase I's in toluene St in toluene 10% in toluene



Alignment of nematic LCs by silylated substrates, under various treatment conditions TABLE V

50 50 50 50 50 50	O'SE Jiod	4 I 592 01 4 I	17 in tolucne sapor phase sapor phase sulution	EOINT EOINT EOINT ABBM	86-96 86-96	Fluka 92360 Fluka 92360 Fluka 92360	I. CHLOROSILANES Monochlorosilanes Trimethyl
\$ 50 50	boil. 55°C boil. 55°C	502 01 A 1	rapor phase solution	TU103 TU103 MBA	86-96 86-96 86-96	Fluka 92360	
\$ 50 50	boil. 55°C boil. 55°C	502 01 A 1	rapor phase solution	TU103 TU103 MBA	86-96 86-96 86-96	Fluka 92360	, éuranne
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ç	•••		nonutos	MBBA	8€-9€		
						07200 -41113	
07	dun moor	nim č1	10°7 in toluene tinsed air dried.	TN403	5.8€-6.8€	Fluka 92360	
			cnted 120°C 1 h				
07	room temp.	nim čl	10% in coluenc	TN404	2.85-2.65	Fluka 92360	
			tinsed air dried,				
00	,	31		CC11.12	300 370	0700 -1-13	
07	room remp.	uim Ci		7011172	C.86C.0t	FIRKS 92500	
02	umal moor	nim 21		13	\$ 81-5 91	Eliit 2 02360	
۰.	dus moor	11111 61		, ,	C:0C-C:0C	00070 99911	
70	room temp.	nim 21		83	2.85-2.85	Fluka 92360	
			rinsed air dried,				
			cured 120°C 1 h				
	0Z 0Z	room temp 20 20 com temp 20 20 20 com temp	05 qmər moor nim 21 05 qmər moor nim 21 05 qmər moor nim 21	1200 1200	TM404 1 1 1 1 1 1 1 1 1	1000 1000	Fluka 92360 36.5–38.5 TM404 TM404 TM304 TM

C'H1O-4-CH=N-4-C-N CH10-4-CH=N-4-C-N C'H10-4-CH=N-4-C-N C'H10-4-CH=N-4-C-N

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	C ₄ H ₁₁ -φ-φ-C≅N C ₇ H ₁ -φ-φ-C≅N CH ₁ -φ-φ-C∃N CH ₁ -φ-C+EN-φ-C ₄ H ₄ C ₇ H ₂ O-φ-CH=N-φ-C ₄ H ₄ C ₇ H ₂ O-φ-CH=N-φ-C ₄ H ₄ CH ₂ O-φ-CH=N-φ-C ₄ H ₄ CH ₂ O-φ-N=N-φ-OCH ₃ O C ₇ H ₄ O-φ-N=N-φ-OC ₇ H ₄ O C ₇ H ₄ O-φ-N=N-φ-OC ₇ H ₁₁ O CH ₁ O-φ-CH=N-φ-OO-C-CH ₁ CH ₂ O-φ-CH=N-φ-OO-C-C ₇ H ₅ CH ₄ O-φ-CH=N-φ=C-N CH ₂ O-φ-CH=N-φ=C-N CH ₃ O-φ-CH=N-φ=C-N CH ₄ O-φ-CH=N-φ=C-N	5% in toluene	10 min		87 87 87 87 87 87 87 87 87	#\T
	C;H ₁ ;-φ-φ-C≡N CH ₂ O-φ-CH=N-φ-C ₄ H ₁ C;H ₂ O-φ-CH=N-φ-C ₄ H ₁ C;H ₂ O-φ-CH=N-φ-C ₄ H ₁ C;H ₃ O-φ-CH=N-φ-C ₄ H ₁ CH ₃ O-φ-N=N-φ-OCH ₁ O C;H ₄ O-φ-N=N-φ-OC;H ₁ O C;H ₁ O-φ-N=N-φ-OC;H ₁ O CH ₃ O-φ-CH=N-φ-OO-C-CH ₁ C;H ₃ O-φ-CH=N-φ-OO-C-C;H ₂ C;H ₃ O-φ-CH=N-φ-OO-C-C;H ₃ C;H ₃ O-φ-CH=N-φ-OO-C-C;H ₄ C;H ₃ O-φ-CH=N-φ-OO-C-C;H ₄ C;H ₃ O-φ-CH=N-φ=OO-C-C;H ₄ C;H ₃ O-φ-CH=N-φ=C-N	5% in toluene	10 min		87 87 87 87 87 87 87 87 87 87 87 87	#\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\
	CH ₁ O-φ-CH=N-φ-C ₄ H ₁ C ₁ H ₁ O-φ-CH=N-φ-C ₄ H ₁ C ₁ H ₁ O-φ-CH=N-φ-C ₄ H ₁ C ₁ H ₂ O-φ-CH=N-φ-C ₄ H ₁ CH ₂ O-φ-CH=N-φ-C ₄ H ₁ CH ₂ O-φ-N=N-φ-OC ₁ H ₁ O C ₁ H ₁ O-φ-N=N-φ-OC ₂ H ₁ O C ₁ H ₁ O-φ-CH=N-φ-OO-C-CH ₁ C ₂ H ₂ O-φ-CH=N-φ-OO-C-C ₂ H ₁ C ₃ H ₄ O-φ-CH=N-φ-OO-C-C ₂ H ₂ C ₄ H ₄ O-φ-CH=N-φ=OO-C-C ₂ H ₃ C ₄ H ₄ O-φ-CH=N-φ=OO-C-C ₃ H ₃ C ₄ H ₄ O-φ-CH=N-φ=C-N	5% in toluene	10 min		87 87 87 87 87 87 87 87 87 87 87 87	#\\T #\\T #\\T #\\T #\\T
	C,H,O-φ-CH=N-φ-C,H, C,H,O-φ-CH=N-φ-C,H, C,H,O-φ-CH=N-φ-C,H, CH,O-φ-CH=N-φ-C,H, CH,O-φ-N=N-φ-OCH, O C,H,O-φ-N=N-φ-OC,H, O C,H,O-φ-CH=N-φ-OC-C-CH, C,H,O-φ-CH=N-φ-OO-C-C,H, C,H,O-ф-CH=N-φ-OO-C-C,H, C,H,O-D-CH=N-φ-OO-C-C,H, C,H,O-D-CH=N-φ-OO-C-C,H, C,H,O-D-CH=N-φ-OO-C-C,H, C,H,O-D-CH=N-φ-OO-C-C,H, C,H,O-D-CH=N-φ-OO-C-C,H, C,H,O-D-CH=N-φ-OO-C-C,H, C,H,O-D-CH=N-φ-OO-C-C,H, C,H,O-D-CH=N-φ-OO-C-C,H, C,H,O-D-CH=N-φ-OO-C-C,H, C,H,O-D-CH=N-ф-OO-C-C,H,	5% in toluene	10 min		87 87 87 87 87 87 87 87 87 87 87	#\T #\T #\T T #\T
	C,H,O-&-CH=N-&-C,H, C,H,O-&-CH=N-&-C,H, CH,O-&-N=N-&-OCH, O C;H,O-&-N=N-&-OC;H, O C;H;O-&-N=N-&-OC;H; O CH;O-&-CH=N-&-OO-C-CH, C;H;O-&-CH=N-&-OO-C-C;H, C;H;O-&-CH=N-&-=C-N	5% in toluene	10 min		87 87 87 87 87 87 87 87 87 87	# # # # # # # # # # # # # # # # # # #
	C,H,O-φ-CH=N-φ-C,H, CH,O-φ-N=N-φ-OCH, O C;H,O-φ-N=N-φ-OC;H, O C;H,IO-φ-N=N-φ-OC;H, O CH,O-φ-CH=N-φ-OO-C-CH, CH,O-φ-CH=N-φ-OO-C-C;H, CH,O-φ-CH=N-φ-OO-C-C;H, CH,O-φ-CH=N-φ-OO-C-C;H, CH,O-φ-CH=N-φ-OO-C-C;H, CH,O-φ-CH=N-φ=O	5% in toluene	10 min		87 87 87 87 87 87 87 87 87	# # # # # # # # # # # # # # # # # # #
	CH,O-φ-N=N-φ-OCH, O O C;H ₁ O-φ-N=N-φ-OC;H ₁ O C;H ₁ O-φ-N=N-φ-OC;H ₁₁ O CH,O-φ-CH=N-φ-OO-C-CH, C;H ₁ O-φ-CH=N-φ-OO-C-C;H ₁ C;H ₂ O-φ-CH=N-φ-OO-C-C;H ₂ C;H ₃ O-φ-CH=N-φ-OO-C-C;H ₃ C;H ₃ O-φ-CH=N-φ-OO-C-C;H ₄ C;H ₃ O-φ-CH=N-φ=C-N C;H ₃ O-φ-CH=N-φ=C-N	5% in toluene	10 min		87 87 87 87 87 87 87 87	# #\\T #\\T T #\\T
	O C;H,O-Φ-N=N-Φ-OC;H, O C;H;O-Φ-N=N-Φ-OC;H;; O CH;O-Φ-CH=N-Φ-OO-C-CH; C;H;O-Φ-CH=N-Φ-OO-C-C;H; C;H;O-Φ-CH=N-Φ-OO-C-C;H; CH;O-Φ-CH=N-Φ-OO-C-C;H; CH;O-Φ-CH=N-Φ-OO-C-C;H; CH;O-Φ-CH=N-Φ=C-N C;H;O-Φ-CH=N-Φ=C-N	5% in toluene	10 min		87 87 87 87 87 87 87	#\T #\T #\T T T
	O C;H ₁₁ O-φ-N=N-φ-OC;H ₁₁ O CH,O-φ-CH=N-φ-OO-C-CH, C;H,O-φ-CH=N-φ-OO-C-CH, C;H,O-φ-CH=N-φ-OO-C-C;H; C;H ₁ O-φ-CH=N-φ-OO-C-C;H, CH,O-φ-CH=N-φ=C-N C;H ₂ O-φ-CH=N-φ=C-N	5% in toluene	10 min		87 87 87 87 87	 \\T \\T \\T
	0 CH,0-φ-CH=N-φ-00-C-CH, C,H,0-φ-CH=N-φ-00-C-CH, CH,0-φ-CH=N-φ-00-C-C;H, CH,0-φ-CH=N-φ-00-C-C;H, CH,0-φ-CH=N-φ=C-N C;H,0-φ-CH=N-φ=C-N	5% in toluene	10 min		87 87 87 87 87	 \T \T \T
	CH ₁ O-φ-CH±N-φ-OO-C-CH ₁ C ₂ H ₃ O-φ-CH±N-φ-OO-C-CH ₃ CH ₃ O-φ-CH±N-φ-OO-C-C ₂ H ₃ C ₃ H ₄ O-φ-CH±N-φ-OO-C-C ₂ H ₄ CH ₃ O-φ-CH±N-φ=C-N C ₂ H ₃ O-φ-CH±N-φ=C-N	5% in toluene	10 min		87 87 87 87	 \T \T
	C ₂ H ₃ O-φ-CH=N-φ-OO-C-CH ₃ CH ₃ O-φ-CH=N-φ-OO-C-C ₂ H ₃ C ₃ H ₄ O-φ-CH=N-φ-OO-C-C ₂ H ₄ CH ₃ O-φ-CH=N-φ=C-N C ₂ H ₃ O-φ-CH=N-φ=C-N	5% in toluene	10 min		87 87 87 87	 \T \T
	CH ₃ O-φ-CH=N-φ-OO-C-C ₂ H ₃ C ₃ H ₄ O-φ-CH=N-φ-OO-C-C ₂ H ₄ CH ₃ O-φ-CH=N-φ≡C-N C ₂ H ₃ O-φ-CH=N-φ≡C-N	5% in toluene	10 min		87 87 87	 \T \T
	C₁H₁O-φ-CH=N-φ-OO-C-C₂H₁ CH₁O-φ-CH=N-φ≡C-N C₂H₁O-φ-CH=N-φ≡C-N	5% in toluene	10 min		87 87	 \T
	CH ₃ O-φ-CH=N-φ≡C-N C ₂ H ₃ O-φ-CH=N-φ≡C-N	5% in toluene	10 min		87	11
	$C_2H_3O-\phi-CH=N-\phi\equiv C-N$			•		
					8/	- 11
	C ₄ H ₄ O-φ-CH=N-φ≡C-N				87	Ä
	C.H.D-φ-CH=N-φ≡C-N				87	ï
	C ₆ H ₁₇ O-φ-CH=N-φ≡C-N				87	ï
	$C_{10}H_{21}O-\phi-CH=N-\phi=C-N$				87	ii
	C ₃ H ₁₁ -φ-φ-C≡N				87	Ï
	C ₆ H ₁₃ -φ-φ-C≡N				87	ii
	$C_1H_{13}-\phi-\phi-C \equiv N$				20	•
	$C_7H_{15}-\phi-\phi$ - $C\equiv N$				87	II
					87	il
						ji
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	CH ₁ O- ϕ -N=N- ϕ -OCH ₁				87	ii
	O					
	C ₂ H ₆ O-φ-N=N-φ-OC ₂ H ₆ O				87	11
	C ₃ H ₁₁ O-φ-N=N-φ-OC ₃ H ₁₁ O				87	1
	CH ₁ O-φ-CH=N-φ-OO-C-CH ₁				87	11/ \(\to \)
	C2H3O-&-CH=N-&-OO-C-CH3				87	/
					87	•
	C ₂ H ₆ O-φ-CH=N-φ-OO-C-C ₂ H ₆				87	II/T
		C ₁ H ₁ ,-φ-¢-C≡N C ₇ H ₁ ,-φ-¢-C≡N C ₇ H ₁ ,-φ-¢-C≡N CH ₁ O-φ-CH=N-φ-C ₄ H ₄ C ₂ H ₃ O-φ-CH=N-φ-C ₄ H ₄ C ₃ H ₃ O-φ-CH=N-φ-C ₄ H ₄ C ₄ H ₄ O-φ-CH=N-φ-C ₄ H ₅ C ₄ H ₄ O-φ-N=N-φ-OCH ₃ O C ₂ H ₄ O-φ-N=N-φ-OC ₇ H ₆ O C ₃ H ₁ O-φ-N=N-φ-OC ₇ H ₁₁ O C ₄ H ₁ O-φ-CH=N-φ-OO-C-CH ₃ C ₇ H ₃ O-φ-CH=N-φ-OO-C-C-CH ₃ C ₇ H ₃ O-φ-CH=N-φ-OO-C-C-C-C-C-C-C-C-C-C-C-C-C-C-C-C-C-	C ₁ H ₁ ,-φ-φ-C≡N C ₁ H ₁ ,-φ-φ-C≡N CH ₁ O ₂ -φ-CE=N CH ₂ O ₃ -φ-C=N CH ₂ O ₄ -CH=N-φ-C ₄ H ₄ C ₂ H ₃ O _{-φ} -CH=N-φ-C ₄ H ₄ C ₃ H ₄ O _{-φ} -CH=N-φ-C ₄ H ₄ C ₄ H ₄ O _{-φ} -N=N-φ-OCH ₃ O C ₃ H ₄ O _{-φ} -N=N-φ-OC;H ₄ O C ₃ H ₄ O _{-φ} -N=N-φ-OC;H ₄ O C ₄ H ₄ O _{-φ} -N=N-φ-OC;H ₄ O CH ₁ O _{-φ} -CH=N-φ-OO-C-CH ₃ CH ₃ O _{-φ} -CH=N-φ-OO-C-CH ₃ CH ₃ O _{-φ} -CH=N-φ-OO-C-CH ₃ CH ₃ O _{-φ} -CH=N-φ-OO-C-CH ₃	C,H ₁ ,-φ-φ-C≡N C,H ₁ ,-φ-φ-C≡N CH,O-φ-CH=N-φ-C,H ₁ C,H,O-φ-CH=N-φ-C,H ₁ C,H,O-φ-CH=N-φ-C,H ₁ C,H,O-φ-CH=N-φ-C,H ₁ CH,O-φ-N=N-φ-OCH ₂ O C,H ₁ O-φ-N=N-φ-OC;H ₁ O C,H ₁ O-φ-N=N-φ-OC;H ₁ O C,H ₁ O-φ-N=N-φ-OC;H ₁ O C,H ₁ O-φ-CH=N-φ-OC-C-CH ₂ C,H ₂ O-φ-CH=N-φ-OO-C-CH ₃ C,H ₃ O-φ-CH=N-φ-OO-C-C,H ₃	C ₁ H ₁ ,-φ-φ-ŒN C ₁ H ₁ ,-φ-φ-ŒN CH ₁ O ₋ φ-φ-C ₄ H ₀ C ₂ H ₂ O ₋ φ-CH=N-φ-C ₄ H ₁ C ₃ H ₃ O ₋ φ-CH=N-φ-C ₄ H ₁ C ₄ H ₄ O ₋ φ-CH=N-φ-C ₄ H ₁ CH ₃ O ₋ φ-N=N-φ-OCH ₃ O C ₃ H ₄ O ₋ φ-N=N-φ-OC ₃ H ₁ O C ₃ H ₁₁ O ₋ φ-N=N-φ-OC ₃ H ₁₁ O CH ₃ O ₋ φ-CH=N-φ-OO-C-CH ₃ C ₃ H ₃ O ₋ φ-CH=N-φ-OO-C-CH ₃ CH ₃ O ₋ φ-CH=N-φ-OO-C-CH ₃	$\begin{array}{cccccccccccccccccccccccccccccccccccc$

TABLE V (continued)

Trichlorosilanes								
Methyl	Fluka 69450	20.3	TN103	vapor phase		boil. 65°C	20	×
Ethyl ²	Fluka 04970	20.5	TNI03	vapor phase	l h	boil. 98°C	20	×
Phenyi ²	Fluka 79230		TNI03	vapor phase	10 sec	boil, 98°C	20	×
Phenyl	Fluka 79230		TN103	spin coated		room temp.	20	×
Phenyl	?		CH ₃ O- ϕ -CH=N- ϕ -C-N	toluene	10 min	room temp.	87	li .
rnenyi			C ₂ H ₃ O-φ-CH=N-φ-C-N	totaciic			87	ii
			C4H4O-6-CH=N-6-C-N				87	ii
			C4H11O-6-CH=N-6-C-N				87	ii
			C ₁ H ₁ ,O-φ-CH=N-φ-C-N				87	ii
							87	ü
			C ₁₀ H ₂₁ O- ϕ -CH=N- ϕ -C-N				87	ï
			C ₄ H ₁₁ -φ-φ-C-N				87	ii
			C ₄ H ₁₃ -φ-φ-C-N				87	ï
			C1H15-φ-φ-C-N				87	ï
			CH ₃ O-φ-CH=N-φ-C ₄ H ₃				87	11/1L
			C ₂ H ₃ O-φ-CH=N-φ-C ₄ H ₉				87	11/1
			C1H1O-6-CH=N-6-C4H1				87	11
			C ₄ H ₉ O-φ-CH=N-φ-C ₄ H ₉				87	1
			CH ₃ O-φ-N=N-φ-OCH ₃ O				87	И
			C2H4O-4-N=N-4-OC2H4				87	11/1
			O C3H11O-ø-N=N-ø-OC3H11				87	1
			0				•	
			CH ₂ O- ϕ -CH=N- ϕ -OO-C-CH ₂				87	IIVT
			$C_2H_3O-\phi-CH=N-\phi-OO-C-CH_3$				87	II/T
			CH ₃ O-φ-CH=N-φ-OO-C-C ₂ H ₃				87	•
			C2H4O-6-CH=N-6-OO-C-C2H4				87	•
Vinyl	Nobel		TN103	spin coated	•••	room temp.	20	on rubbed subst.
Octadecyl (or stearyl)	Fluka		E7	1% in toluene	l h	room temp.	20	T
				curing 130°C				
				20 min (fresh				
				solutions)				.,
			1132					×
			1275					nearly 1
			E8	0.5% in isopropyl	45 min	room temp.		1
				alcohol curing 103°C 1 h				
			404					1
			1132					•
			1275				20	nearly ±
Stearyl			CH ₁ O-φ-CH=N-φ-C-N	5% in toluene	10 min	room temp.	87	И
occal yi			C ₁ H ₄ O-φ-CH=N-φ-C-N			-	87	11
			C4H4O-\$\phi\$-CH=N-\$\phi\$-C-N				87	JI
			C ₄ H ₁₃ O-φ-CH=N-φ-C-N				87	ll .
			C411130-4-C11211-4-C-11					

	.ngilA 	Ref. 87	O°.qm∍T	əmiT	Treatment	Liquid ctystal		ا ر۔10	Supplier	Silane
	H H H	78 78 78 78				C+H ¹ - Q -Q-C-N C ⁴ H ¹¹ -Q-Q-C-N C ⁴ H ¹¹ -Q-Q-C-N C ¹⁰ H ³ 1O-Q-CH-N-Q-C-N				
	 	78 78 78 78				CH'0-や-N=N-や-OCH' C'H'0-や-CH=N-や-C'H' C'H'0-や-CH=N-や-C'H' C'H'0-や-CH=N-や-C'H' CH'0-や-CH=N-や-C'H'				
•	T/II	48				0 C³H'O-や-N=N-や-OC³H' O				
J.	T/∥ T	78 78				CH10-4-CH=N-4-00-C-CH1 O C1H110-4-N=N-4-OC1H11				
COGNARD	● T/● T/	78 78 78				C1H'O-P-CH=N-P-00-C-C'H' CH'O-P-CH=N-P-00-C-C'H' C'H'O-P-CH=N-P-00-C-C-H'				
Ð	X (on rubbed subst.)	02	Doil. (110°C)	41	vapor phase	TNI03		54	Fluka 40120	Dimethyl DES
	× nearly × ×	∠! ≯8	room temp.	nim č1	1938W ni %2.0	MBBA ABBA	(\$8)	53	Shin Etsu chem	SMT IqdisM
	×	70 70	boil. (110°C) room temp.	41	esang rogav betago nige	PBA TV103 TV103	(55	27 25 (glas	Fluka 04960 Union Carbide A-171	Ethyl TES Vinyl TMS
	rubbed subst.		room temp.	•••	bateoa niqe	FOINT		\$7	D)n. Nobel VTMO	ant time
	X ● T	78 78	room temp. room temp.	nim čl nim čl	1918-W 6 ni %2.0 1918-W 6 ni %2.0	WBBY SCB	(84) (84)	77	Shin Ersu chem Shin Ersu chem	Pentyl TES
	×	†8 †8 †8	room temp.	nim &! nim &!	0.5% in 6 water 1\1915 w 8 in 6/29 formulann	PBA SCB MBBA	(\$8) (\$8)	32	Shin Etsu chem Toray	Phenyl TES
	×	70 84	room temp.	•••	possessings	MBBA PBA TN103		2.66	Silicone Union Carbide A-172	yxodism-&-lyniV
	×	20	room temp.	•••	barsos niga	TNIO3	Pyrex)		Union Carbide A-151	TES Vinyl TES
	× ×	LL LL	гоот 1етр.	nim č	0.1% in water, dryed at room temp, cured in	МВВУ + 10% СИ СВОУ МВВУ			Dow Corning XZ-2024	Meihylaminopropyl TMS (MAM)
									••	-
	×	84	ems; m003	nim č1	198w ni %2.0	vadN	1707	UL	7.000 MA: O'	
	X T X	50	room temp.	41	onsulot ni %!	MBBA PBA SCB TMI03	(58)	52 (413-	Dow Corning XZ-2024	Outdoor 1:
	T T	70 70	toom temp.	 nim č	baseos nige	EDINT FOINT	(28) (22) (28) (22)	isla) čč	Wacker A 1100 Dyn. Nobel AMEO	23T lyqo1qonimA-£
	T	07 07	room temp.	mm c	0.1% in toluene + sol. rinse	+0+ LN 103	(28) (ss	salg) čč	Dyn. Nobel AMEO	
∢	T X T	\$8 07	гоош зешь:	nim čl	1916W ni %C.O	808 1133 E7	(28) (28) (28) (28)		Dyn, Nobel AMEO Dyn, Nobel AMEO Tokyo Kasei	
ALIGNMENT	 after	50 84 84	гоот (етр.	•••	baseon nigs	ABBM BAB 101NT	(28)	SE	Kogyo Co. Kogyo Co. Dyn. Nobel AMMO	SMT lyqonqonimA-£
MENT	Britanh ◆	20 20	room temp.	 nim č	spin coated 0.1% in toluene	1/103 E1	(28) (28)	SE SE	Dyn. Nobel AMMO Dyn. Nobel AMMO	
OF	, ×	50 50			+ sol. rinse	E7 404			Dyn. Nobel AMMO Dyn. Nobel AMMO	
LIQL	×	° 50	Boil. 260°C	nim O£	vapor phase	1132 TN103	(84)	8€-9€	Dyn. Nobel AMMO Wacker GF 91	-E-lydisonimA-S-N ZMT lyqoiqonimA
ло с	X T T	20 20 20	room temp. room temp. room temp.	 nim č	spin coated spin coated 0.1% in toluene	LN103 E1 LN103	(28) (28) (28)	8€-9€ 8€-9€	Dyn, Nobel DAMO Wacker GF 91	
LIQUID CRYSTALS	×	0Z 0Z			+ sol, rinse	E7 404				
ALS	× ×	78 78	room temp.	nim &! nim &!	moon is grityib	WBBA SCB 1132	(\$4)	9£	Shin Etsu chem Shin Etsu chem	-E-lydsonimA-S-N ZMT lyqorqonimA
	wols T	50 50 84	room temp.	nim č! 	temp. 130°C curing for 20 min spin coated	8A9 E0107	(28)	9E 9E	Shin Elsu chem Dyn. Nobel DAMO	-6-lydisonimA-S-W
	L after heating	07 07	room temp.	•••	batsoa niqe batsoa niqe	1/103 E7	(78)	9€	Dyn. Nobel DAMO Dyn. Nobel TRIAMO	SMT lygorgonimA SMT onimsirT
	baddur Isdus	0Z	тоот сетр.	ním č	0.1% in toluene + sol. rinse	EOINI			OMAIRT	
	baddur Jedus	07	toom temp.	•••	0.1% in toluene + sol. rinse	101			ОМАІЯТ	
29	tnppeq	20	room temp.		0.1% in toluene	£7			OMAIRT	

Silane	Supplier	10 ⁻³ J	m-2	Liquid crystal	Treatment	Time	Temp. °C	Ref.	Align.
	TRIAMO			1132	0.1% in toluene + sol. rinse	•••	room temp.	20	rubbed subst.
	TRIAMO			E7	spin coated		room temp.	20	•
3-Chloropropyl TMS	Dyn. Nobel CPTMO	40.5	(82)	TNI03	spin coated	•••	room temp.	20	rubbed subst.
3-Chloropropyl TES	Dyn. Nobel CPTEO	40.5	(82)	TN103	spin coated	•••	room temp.	20	rubbed subst.
3-Głyceryloxypropył TMS	Dyn. Nobel GLYMO	38-42	(82)	TN103	spin coated		room temp.	20	rubbed subst.
3-Glycerylpropyl TMS	Shin Etsu chem	38.5	(84)	SCB MBBA PBA	1.5% in water	15 min	room temp.	84	× ×
3-Morpholinopropyl TMS	Shin Etsu chem	42	(84)	SCB MBBA BAB	0.5% in water	15 min	room temp.	84	× × ×
3-Methacryloxypropyl TMS	Shin Etsu chem	28/31 28	(84)	SCB MBBA PBA	1% in water/ I methanol			84	(X) X X
Hexamethyldisilarane	Fluka 52619			TN103	vapor phase	l h	boil. 124°C	20	×
	Fluka 52619			TN103	spin coated	•••	room temp.	20	×
Imidazolin	Dyn. Nobel IMEO			TN103	0.1% in toluene + sol. rinse	5 min	room temp.	20	rubbed subst.
				404 E7				20 20	rubbed subst. rubbed
				1132				20	subst. rubbed
				1132			*.		subst.
Imidazol TES	Dyn. Nobel IMEO			TN103	spin coated	•••	room temp.	20	⊥ after heating ●
				E7	spin coated		room temp.		
V-Phenyl-3- aminopropyl TMS	Shin Etsu chem	42	(84)	5CB	3% in 5 water/ I methanol	15 min	room temp.	84 84	nearly ×
	Shin Eisu chem	MBBA		MBBA				84	×
-Mercaptopropyl TMS	Shin Etsu chem	MBBA 41-42	(84)	PBA SCB	3% in 7 water/ 1 methanol	15 min	room temp.	84	ê
		41	(82)	MBBA	· memano.			84	•
		41	(82)	PBA				84	×
Polyamino TMS	Toray	26 .	(84)	5CB	0.5% in water	15 min	room temp.	84	nearly X
								•	
	Silicone	26 26	(84)	MBBA PBA					x .
N,N-Dimethyl-N-	Dow Corning	26	(84) (84)	MBBA	0.1% in water	5 min	room temp.	(77)	Τ
Octadecyl-3-amino- propyl TMS-	XZ-2-2300 Dow Corning	26	(84)	СВОА			•	•	1
chloride (DMOAP)	XZ-2-2300 Dow Corning	26	(84)	MBBA + esters	0.5% in water	10 min	75°C		1
	XZ-2-2300 Dow Corning	26	(84)	MBBA/EBBA					Τ
	XZ-2-2300 Dow Corning XZ-2-2300	26	(84)	n-azoxy LC mixt.	•				Τ
OMOAP	72-2-2500 Toray	23	(84)	5CB	water	15 min	room temp.	(84)	1

^{*}Concerning the difference between vapor and solution coated substrates, see also Ref. 88. T.E.S. = triethoxysilane; T.M.S. = trimethoxysilane. ¹ γ_c from Ref. 82 except as otherwise indicated. ² These silanes hydrolyze strongly, ¹ || and × are often not distinguished in the literature. In our observations only rubbed surfaces either before or after silane treatment lead to || alignment for these silanes which do not induce homeotropy.

⊥ = homeotropic alignment (perpendicular to glass surface): || = alignment, uniform, parallel to glass surface; ● = molecular alignment too difficult to distinguish between perpendicular and parallel alignments and to obtain consistent experimental results: ||/⊥ = molecular alignment changed near the phase transition temperature from the parallel alignment at lower temperatures to the perpendicular alignment at higher temperatures; × = parallel nonuniform.

hazardous. From Ref. 88 it seems that the surface configuration of silane films methyldichlorosilane and phenyltrichlorosilane, in EBBA-MBBA has been preclude glass frit sealing. Treatment of the assembled cell by the organic vapor could be used, but the toxicity of chlorosilanes renders the process deposited from vapor and solution phases is different. Dissolution of diused to promote homeotropic and homogeneous alignment, respectively, but the process is impractical as the conductivity is increased and $T_{\rm c}$ decreased

1.5 Tilted alignment

In evaluation studies of alignment effects, classification of LC alignment into fact, the nematic director of LC molecules generally makes an angle ϕ with the plays in order to obtain a rotation of all the LC molecules in the same direction on the application of an electric field. The tilt supresses the formation of difnonhomeotropic alignment is referred to as parallel (or homogeneous). In substrate surface. This "tilt angle" is required in practical twisted nematic disusing walls between domains with reverse twist. A careful choice of tilt on homeotropic or parallel is an oversimplification, as often, in the literature, each electrode leads to a better optical appearance.

he LC director in the bulk. Literature data show some variations in tilt angle values which arise either from the precision of the measurements or from dif-In the neighborhood of a particular surface, tilt angles are distributed cussed 94 and some indicative data are collected in Table VI. The tilt angle on The rate of evaporation, 46 the pressure, 95 boat temperature, and any subsequent thermal treatment 96 also changes ϕ by several degrees. The variation of ϕ with evaporation rate has been used to monitor the tilt angle of different LC composition. 97 The nature of the LC material, 93,95 its purity, 93 and—as shown in the case of cyanobiphenyls—the chain length, 85 also modify tilt values. We have observed wide variations of the tilt angle values in various cells where measurements have been made just after filling and subsequently after a Despite experimental discrepancies, in general, rubbed layers give low tilt anaround a mean value. 33 Experimental figures refer to the mean orientation of ferent experimental conditions. Accuracy of measurement has been dis-SiOx evaporated layers shows slight variations with the substrate. Layer thicknesses in the range 50-100 Å cause variations of ϕ from 16° to 22° 95 period of stabilization: measurements on 11 cells have shown that, for ROTN 200, on obliquely evaporated SiO_x (heta : 85°), the tilt angle values vary from 13° to 23°; however, after 500 hr, the different cells gave $\phi = 17.5^{\circ} \pm 0.5^{\circ}$. gles and rubbed polymers produce larger tilt angles (1°-5°) than rubbed inorganic surfaces (0°-2°), although 0° tilt angle has been measured on rubbed polyvinyl alcohol for a wide range of LC mixtures 95 and $\phi=1^{\circ}$ has been measured for an ester mixture aligned by polyamide-imide layers deposited on

ALIGNMENT OF LIQUID CRYSTALS

33

Indicative values of nematic director tilt angles on some typical aligning layers

Substrate	CC	Angle (°)	Method	Reference
Evaporated: SiO _x 63°	٠.	3%	magneto canacit	(102)
SiO _r 85°	SCB	27	moving isogvre	(96)
	ecb	29	moving isogvre	(96)
	eCB	29.4 ± 0.7	Δn	(110)
	7CB	31	moving isogyre	(96)
	7CB	24.1	magneto capacit.	(94)
	8CB	34	moving isogyre	(96)
	SOCB	34	•	
	70CB	37		
	90CB	39		
	60CB	38		
	80CB	14	moving isogyre	(96)
SiO _x 85°	E8	25-27 (25	magneto capacit.	(5)
		after 300 h)		
SiO, 82°	E7	25.3	magneto capacit.	(94)
	MBBA	23.3-26	magneto capacit.	(94)
	TN103	21.8	magneto capacit.	(62)
	TN200	15-22 (17	•	
		after 910 h)		(20)
	TN200	22.2		(62)
	TN403	22.7		
i	E7	25.4		
SiO, 82°	E7	25.3	magneto capacit.	(62)
	E7. TN200	25 - 0	capacitance	(104)
100 A/0 - 30 A	1			
SiO _x 60°/85° 100 Å/0 – 10 Å	E7	0 - 25		(103)
	(5CB/60CB)	0 - 33		(111)
SiO _x 85 → 80°	nSB	30 45	Canacitance	(111)
84 - 74°	nSB	15 - 25	magneto canacit	()
76 - 72°	6		magneto capacit.	£ (£
Rubbed:			magneto capacit.	(101)
In ₂ O ₃	E3	0.05	magneto null	(63)
Glass	SB	0.5-5	Canacitance	(2)
PVA	103.200.	0	magneto canacit	(41)
	403. E7			(2)
In ₂ O ₃	E7	2.15 ± 0.1	magneto capacit.	(112)
NMAP silane	E	0	magneto null.	(63)
Rhodiakermid	E3	0.95	magneto null.	(63)
				/ /

pentylcyanobiphenyl on rubbed glass as well as for a biphenyl mixture on rubbed SiOx layers evaporated at 60°C indicence. 95 SiO2 layers rubbed with SiO₂ rubbed with diamond paste.⁸⁹ A tilt angle of 2° has been measured for diamond paste align ester mixtures parallel to the substrate ($\phi=0^\circ$). ⁸⁹

Evaporation of metals or oxides obliquely to the substrate leads to layers which align LCs differently depending on the angle heta of the evaporation beam

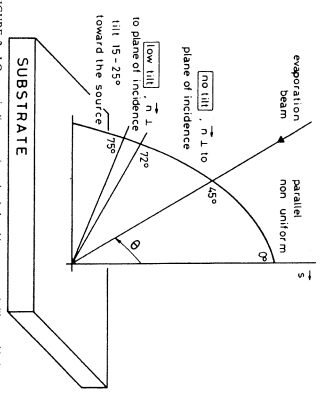


FIGURE 2 LC nematic director orientation induced by evaporated silicon oxide (or magnesium fluoride). Depending upon the angle of incidence of the evaporation beam different orientations are observed.

with respect to the normal to the surface⁴⁴ (Figure 2). If $\theta < 45^\circ$, parallel but nonuniform alignment is observed, but when $45^\circ < \theta < 75^\circ$ LC molecules align in a direction perpendicular to the incident beam with a tilt angle very close to zero. Tangential evaporation at $\theta > 75^\circ$ results in the nematic director being oriented towards the source at an angle of approximately 15°, increasing to 25° as θ increases from 75° to 88°. ^{48,102} When θ is 72°–75° the LC molecules lie parallel to the substrate plane, with the director axis perpendicular to the plane of incidence of the evaporation beam, at a low tilt angle (3°–9°). ¹⁰² These values have been widely confirmed by numerous authors and repeatedly obtained in industrial practice for all of the practical LC mixtures.

Tangentially evaporated MgF₂ gives similar results but with smaller tilt angles than SiO_x for the same angle of evaporation. ^{13b} Oblique evaporation of CaF₂ is reported ¹² to produce homeotropic alignment ($\phi = 90^{\circ}$) when $\theta < 45^{\circ}$, $\phi = 75^{\circ}$ for $45 < \theta < 70^{\circ}$ and $\phi = 60^{\circ}$ for tangentially evaporated layers ($\theta > 75^{\circ}$). Surfactants, some silanes and polymers produce homeotropic alignment. Rubbing and tangential evaporation of oxides give low tilt angles while oblique evaporation induces a 25° tilt angle. When other tilt angles are desired, any combination of high and low-tilt-angle-producing

ent thickness at the same angle of incidence θ . Deposition of a very thin (less may be lowered by rubbing" or by crossed evaporation of two layers of differa rotation of 90°, and 100 Å evaporation of SiO_x at 80°, low tilt angles (5°) are than 5 Å) layer of SiOx, evaporated at an angle of 6° from the substrate methods can be employed. The tilt angle of a tangentially evaporated layer allows tilt angles to be controlled from 0° to 30°. Variation of ϕ from 0 to 45° obtained with the LCs E7 or ROTN 200. Variation of the layer's thickness initial evaporation of a 50 Å SiO, layer at an incidence angle of 5°, followed by plate, gives a tilt angle of 0° to 6° . ¹⁰² By employing the reverse order ¹⁰⁴ i.e., an $(\theta = 84^{\circ})$, on to a layer deposited at 60° incidence, after 90° rotation of the sources. 105 Electron gun deposition of an initial layer, 350 Å thick, at 6° incilayer on to 70° evaporated SiO_x gives a tilt angle of 15° 106 ing on the surfactant chain length, 456 while plasma deposition of an 8 Å Teflon with long chain alcohol allows tilt angles to be varied from 75° to 90° dependduced by monitoring the thickness of the second layer. SAIBE SiO2 covered the substrate, gives a 6° tilt angle. 102 Variations of ϕ from 11° to 3° can be prodence, followed by a 20-50 Å thick layer deposited at 30° without rotation of for E₃ has been obtained by simultaneous evaporation of SiO from two

A SiO_x layer evaporated at a low angle, which would normally result in a tilt angle of 25°, when treated with a homeotropic aligning layer of lecithin, orient the LC at the complementary angle of $\phi = 65^\circ$ (Figure 3), as has been observed when this layer is treated with octadecylammonium bromide, 40 or deposited on glass cleaned with sulfochromic acid. 108 Evaporation of two dissimilar materials, such as gold and indium oxide, on adjacent sides of the grooves of a wavy surface gives a LC alignment slightly tilted from the normal for LC mixtures containing a homeotropically aligning surfactant. 107

Orientation of thin LC layers may be obtained with the nematic director oriented in any direction from $\phi = 0$ to 25°-35° and 60° to 90° by a suitable aligning layer (Diagram 1).

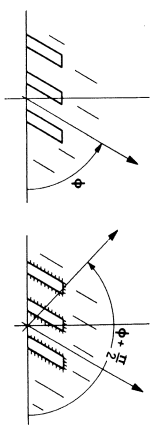


FIGURE 3 Where an inorganic surface produces an average orientation of the LC molecules at an angle φ, treatment by an homeotropic aligning surfactant leads to an orientation at the complementary angle.

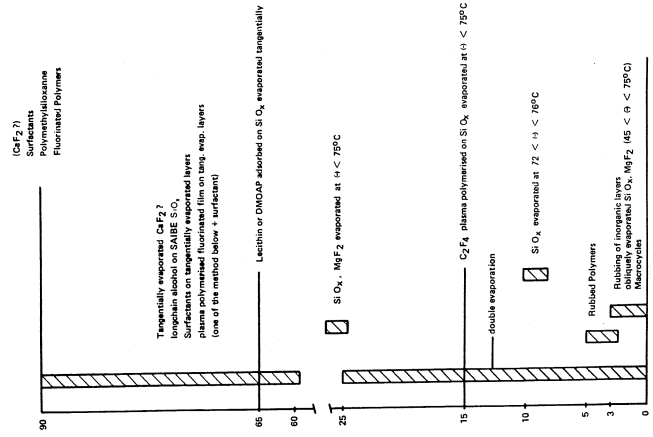


DIAGRAM 1 Tilt angle of the LC nematic director produced by different surface treatment.

ALIGNMENT OF LIQUID CRYSTALS

II ALIGNMENT MECHANISMS OF NEMATIC LIQUID CRYSTALS AND THEIR MIXTURES

Evaluating the various alignment methods reviewed in § I aided greatly in clarifying some general guidelines for LC-surface interactions.

The difficulties encountered, while attempting to obtain meaningful experimental data, due to ill-defined substrates, LC decomposition, and surface chemistry misconception have hindered progress in the understanding of the subject. LC molecules are composed of an aromatic core, an alkyl chain and a polar group, thus giving rise to complex dispersive polar and hydrophobic interactions.

II.1 Physicochemical interactions

II.1.1 The Friedel-Creagh-Kmetz rule

An old, but neglected study of the alignment of LCs (mainly azoxy compounds) on freshly cleaved crystals²³ suggested that the orientation of the LC molecules was determined by their physiocochemical interactions: of 80 studies of LC alignment on crystal surfaces, 77 showed parallel orientation. If, however, a hole was drilled in the solid substrate, homeotropic alignment ensued in the suspended film. This led to the conclusion that LC films aligned parallel to a solid substrate if any interaction occurred and homeotropically in the absence of interactions.¹⁰

Solid-liquid surface interactions are phenomenologically described by the surface tensions of the solid substrate, γ_8 and of the liquid γ_L . γ_L is a macroscopic description of the strenght of the LC-LC interaction while γ_8 describes the solid surface excess energy.

Thus, the above observations may be stated as⁶

$$\gamma_S < \gamma_{LC}$$
 homeotropic alignment $\gamma_S > \gamma_{LC}$ parallel alignment

This empirical rule has been widely supported by experimental data, although not all results are in agreement. We will call it the "Friedel-Creagh-Kmetz" (FCK) rule.

II.1.2 Surface tension

II.1.2.1 Liquid crystal surface tension

In order to have thermodynamic significance, the LC surface tension, γ_{LV} , should be measured with the LC in equilibrium with its vapor. Most of the available data are relevant to an air-LC interface, γ_{LA} . Although the precision

equilibrium conditions and/or atmospheric contamination of the LC (for the $\rm J~m^{-2}$, comparison of the published values, (Table VII), shows a dispersion, of the different methods used to measure the LC surface tension should be 10⁻⁴ influence of different methods see § II.1.10). that may amount to 30% with the same method. This is probably due to non-

order of 25-40 erg/cm cisely, 113 as the LC interface drifts continuously. LC surface tensions are of the The surface tension of MBBA has been found impossible to measure pre-

II.1.2.2 Anisotropy of the surface tension of a LC

molecules were oriented parallel or perpendicular to the surface. cules at the surface, it would be expected to vary according to whether the LC lar potential. As surface tension results from the excess energy of the mole-The existence of a liquid crystalline state implies an anisotropic intermolecu-

axis dipole moment (μ_1) averages out. As a result the LC molecules lie flat at der Waals interactions 118,119 leads to $\gamma_{\parallel}<\gamma_{\perp}$. This calculation is applicable to the LC-air interface (e.g., PAA, Ref. 115). LCs of negative $\Delta \epsilon$ as their long axis dipole moment (μ_{\parallel}) is weak and the off Calculations of the surface tension anisotropy, taking into account only van

retical evaluation of dipole interactions in LCs. ¹¹⁹ The calculated value of $\gamma_{\parallel} - \gamma_{\perp}$ is 7×10^{-3} J m⁻² for PAA (Ref. 116) while $\gamma_{\parallel} - \gamma_{\perp}$ of the order of align normal to the free surface 120; thus $\gamma_{\parallel} < \gamma_{\perp}$, which contradicts the theo- $5 \times 10^{-6} \, \mathrm{J \, m^{-2}}$ (Ref. 75, 120) is obtained from experimental observations (see could be due to the angle between the molecular axis and the diple moment. [10 - 7S/6] (Ref. 119)† with the previously mentioned limitations. Table VII). The Kirkwood-Buff approximation leads to $\gamma_{\parallel}-\gamma_{\perp}=\gamma_{0}(S/9)$ For LCs of strong positive $\Delta\epsilon$, experimental observations show that the LCs MBBA has $\mu_{\parallel} = 0.4$ D and is slightly tilted 114 at the free surface, which

calculated from theory. Further work seems to be required to ascertain the small to explain the observed tilted alignment of MBBA on glasses onto which real order of magnitude and its relation to various LC classes. $(\gamma s \sim 26 \times 10^{-3} \, \mathrm{J \, m^{-2}})$. Such an effect would require $\Delta \gamma \ge 8 \times 10^{-3} \, \mathrm{J \, m^{-2}}$ as fatty derivatives of medium chain lenght (C6-C10) have been adsorbed tilted orientation of the nematic director. Experimental values are much too surface tension, γ_s , has an intermediate value $\gamma_{\parallel} < \gamma_s < \gamma_{\perp}$, leading to a It has been suggested' that a situation could exist where the solid substrate

II.1.2.3 The surface tension of solids

excess surface energy from the energy of the broken bonds necessary to create The surface tension of a solid cannot be measured directly. Calculation of the

a surface, give values of 0.5 to 5 J m^{-2 121}; these values refer to the solid-

vacuum interface, γ8.

constituants strongly adsorb onto the surface, lowering the surface tension from the uncontaminated value of the "surface pressure" π_0 . As soon as the solid surface is brought into contact with air, atmospheric

The solid-air interface has thus a surface energy,

$$\gamma_{\rm SA} \approx \gamma_{\rm S}^{\rm g} - \pi_0$$

where π_0 is approximately 0.1 J m⁻² 128

layer of water, with $\gamma_{SA} = 7 \times 10^{-2} \text{ J m}^{-2.117}$ been suggested that oxide surfaces should be considered as covered by a thin Due to the fact that the solid surface chemical state is poorly defined, it has

sorbed water causes TiO2 surfaces to become hydrophobic. 123 750°C, the silica surface become irreversibly hydrophobic. 122 Irreversibly abbridges. Above 400°C, the oxygen bonding becomes irreversible, and above sation of surface hydroxy groups between 170°C and 400°C, to form oxygen physically absorbed water at around 150°C and following this will be condenproperties of a solid surface. In the case of silica layers, heating will desorb the Cleaning procedures, as well as further thermal treatment, will modify the

which are not suitable for glass cells. inorganic substrates supposes thermal treatment at very high temperatures, In general, hydrophobic oxide surfaces consist of areas incapable of specific molecular interactions. ¹²³ Complete removal of the specific interactions on

Tin oxide thin films, formed by CVD are hydrophobic; acid cleaning renders them hydrophilic.²⁰

however, γ_{SA} is greater than 4×10^{-2} J m⁻² during display sealing) and cleaning procedures. For inorganic solid surfaces, preparation conditions, subsequent thermal treatment, (which is unavoidable predict. The same oxide may show different surface energies, depending on Even in the case of a uniform solid surface, its surface energy is difficult to

the situation is not worse than that of surface chemistry. the absorbed LC film. It is often claimed that LC alignment is not understood; $(\gamma_{\rm SV^0})$, at the saturated vapor pressure. In this case π_0 would be the pressure of measure the surface tension of the solid in equilibrium with the LC vapor To reliably establish the validity of the FCK rule it would be necessary to

II.1.3 Solid-liquid crystal contact angle

II.1.3.1 The contact angle

drop. The angle $heta_{SL}$ between the edge of the drop and the surface is related to YLV, YSV and the interfacial energy, YSL, by the expression When a liquid is deposited on a surface it often happens that it remains as a

$$\gamma_{SV} \cos \theta_{SL} = \gamma_{SV^{\circ}} - \gamma_{SL}. \tag{3.1}$$

[†] Reference 113 suggests that after correction of errors in the evaluation of the integral, $\gamma_1 = \gamma_0 (S/3)(5 - 19/32S)$.

Surface tension ye of some nematic LCs* TABLE VII

sArsmaFl		borizeM	.ħeÆ	2- ML ^{E-} 01	ר' כי	×
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Fifect of impurities is slight (38)	•	***	/11	86	A88M	HIFF'S BASE
MBBA aligns nearly ⊥ at the free surface (114 b) ⇒	•	IA	bil	38 ∓ 3 (53 ₀ C)		
TL /L</td <td></td> <td>111</td> <td>38</td> <td>35.8</td> <td></td> <td></td>		111	38	35.8		
\/ = 3\tau \\ = 3e \\ (2)	•	Ш	911	32'3 (52 ₀ C)		
		۸	SII	6.36		
$(211) 6 = {}^{d}\mathcal{L} 82 = {}^{p}\mathcal{L}$	•	н	7.6	34		
$\chi_{d} = 33 \qquad \gamma_{p} = 1 \qquad (37)$		ì	34	34		
corr - uncorrected	İ))	9	32 - 34		
γ _L = 28.8 corr. Harkins and Jordan (1)		11	ı	32.6		
	1	1	S	30		
$(12) \qquad S^{*}\iota = {}^{d} \mathcal{L} \qquad \mathfrak{p} \mathfrak{E} = {}^{p} \mathcal{L}$	•	H	7E	36.85	\A88M 08	
					50 BBC∀	
← anigne # to the free surface	•	11	PZII	4.0 ± 8.6€	AA9	SYXO
$/\!\!/ L < ^{T} L$		шл	9 411	(30 ₀ CI) 6E	(150 ₀ C)	
		Λ	Sii	6.86		
(013) 377 = pro 020 (7 7 p		НΛ	138	\$.0 ± 8.8€		
(811) 8.44 = 17 8.75 = (also) 1/b 7	•	IΛ	9 111	38 ∓ 4		
		1117	2711	6.7€		
	- 1	111	911	37 (120°C)	i !	

4	IIV mentenna animattenna tidait IV masiliana Vanta						
	,						
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	,						
κ.			111	SO	30.6 ± 0.2 (22°C)	83	
AL			111	SO	(20°C) (22°C)	£3	
/ST	_		ŀ	EII	39'3. (41 ₀ C)	8 CB	
CR.	υ _τ = 30	•	1	ειι	57.9 (20°C)		
OF LIQUID CRYSTALS	(Insmngils pigottosmort tot besteatt sets19)	•	ł	EII	28.1 (31.5°C)		
ΔΩ				70	30		
Ĭ	•		1 111	35 50	32 ± 0.5		
OF	T L < #L		11	1/8	3.45		
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ME	<u> </u>						
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ALI.			111	50	27.5 ± 0.3 (22°C)	EOI NTOR	
`			ı	34	72	HCPB	0 < ∍ ♡
			1	34	24.5	aggM	0 > 3 △ A3T23
			111	38	7.36	BECS	STILBENE
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			11	6 TII	8.62		
			۸	511	2.62	(128 _o C)	
			IIIA	9 (11)	1,92	949	
		- 1	,,,,,	1 7	1		

^{*} Methods: I, wilhemy plate; II, de Nouy ring; III, hanging drop; IV, contact angle; V, capillary; VI, light scattering spectrum. VIII, radii of curvature; VIII, maximum bubble pressure.

At equilibrium it should not matter whether the drop is deposited on the surface (advancing angle) or whether it is formed by pulling out a film of the liquid (receding angle). ¹²⁴ On a LC drop, in practice, variations in θ_{SL} as great as 50° may be observed, and experimentalists generally choose the advancing angle. When studying hydrophobicity, the receeding angle is a better characterization of the surface.

The existence of hysteresis has been attributed to contamination of either the liquid or the solid, rough surfaces or absorbed surface film immobility. 121

II.1.3.2 The contact of a liquid crystal with a solid

Whatever the method used to measure the contact angle of a LC drop on a solid, observations are hampered by the continuous drift of θ_{SL} . On the one hand the absorbed atmospheric constituants have to exchange with the LC molecules; on the other hand the elastic energy of the misaligned layer is weak and the evolution of surface attached disclinations is slow. Four days have been selected as the time necessary for a LC to attain equilibrium with an adjacent surface when making accurate measurements. ¹¹³ Liquid crystal hydrolysis and/or oxidation under the influence of both UV light and oxygen drolysis and/or oxidation under the influence of both UV light and oxygen served to be homeotropically aligned after 10 days, due to the formation of polar decomposition products.

On substrates which induce parallel LC alignment, the viscous flow of the LC tends to determine the direction of alignment, but many defects are formed and they evolve constantly, resulting in constant movement of the LC drop. It seems that reliable values can only be obtained with carefully controlled experimental conditions which has rarely been the case in the published studies.

II.1.4 The critical surface tension of a solid

A useful empirical parameter has been proposed to characterize a solid surface. It has been observed that the contact angle θ_{SL} of homologous alkanes varies linearly with the liquid surface energy. Extropolation to $\cos \theta_{SL} = 1$ led to a critical value γ_C which was characteristic of the solid employed. ¹²⁸

From Eq. (3.1)

$$\gamma_{LV}\cos\theta=\gamma_{SV}-\gamma_{SL}$$

and, therefore

$$\gamma_{\rm C} = \gamma_{\rm SV} - \gamma_{\rm SL}$$

The introduction of a critical surface tension is only valid inasmuch as the values γ_{SL} and π_0 ($\gamma_{SV} = \gamma_{S^0} - \pi_0$) of different liquids are constant. If this is the case for polymers and nonpolar liquids, the use of polar liquids to measure

 $\gamma_{\rm C}$ may cause large variations of the extrapolated value. Therefore, $\gamma_{\rm C}$ values for polar oxide surfaces relate to the nature of the surface and the choice of the solvents. The following observations indicate that in these cases $\gamma_{\rm C}$ is not a useful parameter for the characterization of LC-oxide interactions.

- The critical surface tension of fused silica is $\gamma_C = 78 \times 10^{-3} \text{ J m}^{-2.125}$
- On SiO_x tangentially evaporated films, γ_c has been measured to be approx. 38×10^{-3} J m⁻² after correcting for surface roughness 126 with an anisotropy which depends on whether the edge of the drop is parallel or perpendicular to the evaporation direction.
- Normal evaporation is reported to produce films of $\gamma_{\rm C}=48.7\times10^{-3}$ J m⁻² while obliquely evaporated layers show 48.7 \times 10⁻³ J m⁻² $<\gamma_{\rm C}<55.9\times10^{-3}$ J m⁻² for evaporation angles between 0° and 60°. 127
- On SAIBE etched SiO₂ surfaces, values of $\gamma_{\rm C}$ as low as 24×10^{-3} J m⁻² have been measured^{45a} which is nearly the critical surface tension of water [$\gamma_{\rm C}({\rm H_2O}) = 22 \times 10^{-3}$ J m⁻²].^{45c}
- Atmospheric humidity causes the critical surface tension of glass to vary from between $\gamma_{\rm C}=75\times 10^{-3}\,{\rm J\,m^{-2}}$ for dry glass, to $\gamma_{\rm C}=30\times 10^{-3}\,{\rm J\,m^{-2}}$ at 100% humidity.
- Although water has a γ_C value of 22 \times 10⁻³ J m⁻², LCs ($\gamma_C = 3 \times 10^{-2}$ J m⁻²) do in fact align parallel on water. ¹²⁰

As developed in the following paragraph γ_C is equal to the solid surface tension γ_{SV^o} when the ratios or polar and dispersive interactions of both phases are equal. For this reason the critical surface tension of polymers may be employed to give a qualitative description of polymer–LC interactions. Measurements made on plasma polymerized films ^{34,37} showed acceptable agreement with theory, although some unexpected results were observed, i.e., tolans $(\gamma_{LC} \sim 22 \times 10^{-3} \text{ J m}^{-2})$ were not aligned homeotropically by a sputtered Teflon layer $(\gamma_C \sim 18 \times 10^{-3} \text{ J m}^{-2})$.

Adsorbed surfactants reduce the solid surface tension. Fatty acids or their derivatives (alcohols and amines) exhibit a continuous range of γ_C values with alkyl chain length^{5,75} but reported relations to LC alignment are somewhat conflicting (Figure 4). The same discrepancies are observed with alkyl trimethyl ammonium bromide derivatives. The influence of the surface coverage (or packing density) is of the utmost importance and consideration of γ_C is generally misleading (see paragraph 18).

The FCK rule has generally been quoted 1.6 using ye in place of ys, i.e.,

 $\gamma_{\rm C} < \gamma_{\rm LC} \longrightarrow {\rm homeotropic}$ $\gamma_{\rm C} > \gamma_{\rm LC} \longrightarrow {\rm parallel\ alignment}$



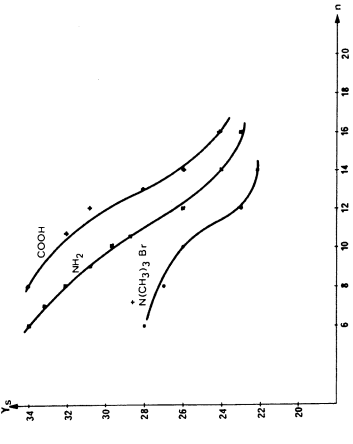


FIGURE 4 Variation of the surface tension of surfactant treated glass surfaces with increasing carbon content of the surfactant alkyl chain. [• from Ref. 75a, × Ref. 5a + Ref. 56].

pared to the expression of § II.1.1, where $\gamma_8 = \gamma_8^0 - \pi_0$, it is evident that the above inequality employs values that are relevant to nonpolar solvents, to describe the behavior of LCs which have to be highly polar for practical and it is, therefore, not surprising that it encountered strong criticism. Comapplication.

11.1.5 The polar and dispersive components of surface tension

Surface interactions are long range forces; either dispersive (van der Waals) forces, dipole-dipole interactions or double layer interactions. 121 These contributions are considered additive and the surface tension $\gamma_i(i=s \text{ or } 1)$ may be written as the sum of the dispersive γ^d_i and polar γ^p_i contributions,

$$\gamma_i = \gamma_i^d + \gamma_i^p.$$

tension: $d_i = \gamma_i^4/\gamma_i$, $p_i = \gamma_i^8/\gamma_i$. It is shown that the critical surface tension γ_c Let di and pi be the ratios of dispersive and polar contributions to the surface

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is $\gamma c = \phi \gamma_{sv}$ where $\phi = \sqrt{d_L d_s} + \sqrt{p_L p_s}$. When $d_i = d_s$, which implies $p_L = p_S$, $\phi = 1$ and $\gamma_C = \gamma_{SV}$.

Thus the empirical method of estimating γ_C may result in a value that agrees well with ys rather than the solid surface tension ysv. 129 Recent work on plasma copolymerized films with different γ^d and γ^p values 37 shows the relation of these considerations to nematic, LC alignment.

11.1.5.1 Polar interactions of LCs and surfaces

ion of the first surface layers, although theoretical understanding of the probem and the experimental data are poor. The effect of surface polarization has been demonstrated in the case of MBBA, 130 for which it was suggested that the MBBA molecules orient with their dipoles towards acid treated glass, but with their butyl chain towards surface cetylammonium bromide monolayers on The interactions between the dipoles of LC molecules and the charged surfaces of oxides or ionic surfactants are certainly very important in the formaIn addition, the surface dipoles of sputter etched indium oxide have in some (but unreproducible) cases been observed 131 to align LCs of negative dielectric anisotropy parallel to the substrate, but positive LCs perpendicular to it. The same behavior is observed with surfaces treated with dicarboxylate 78b chromium complexes, suggesting that only part of the acid group is absorbed onto the substrate. The polar interaction of MBBA with a humid glass substrate, \dagger where $E_s = 4.35 \times 10^9 \text{ V/m}$, have been shown¹³² to be accounted for through the surface electric field, $E_{
m S}$ (Table VIII) as the polar part $W_{
m a}$ of adhesion energy. W_{\blacksquare} is related to the LCs molecular dipole through

 μ effective dipole moment). $W_{a}p = \delta \mu E_{s}$ (δ effective dipole density)

In view of this the existence of a double layer (§ II.1.6) should be considered.

II.1.5.2 Dispersive interactions of LCs and polymers

As already mentioned, nonpolar surfaces offer an important simplification in the theoretical description of liquid-solid interactions as the polar term becomes negligible regardless of the character of the liquid molecules. Most organic compounds as well as Liquid Crystals and polymers have d > p.

Denoting γ^d as the dispersive component of the surface tension, the

[†] This work has been extended to 5CB by these authors in Ref. 133 but the observation of homeotropic alignment of 5CB on glass cast some doubts on their conclusions.

Electrostatic field strength \mathcal{E}_8 and dispersion energies γ_0^4 for some inorganic substrates (from Ref. 132)

HIN STRYL

Author Rustice (bare) AlpOy-SiOy coated CaFy SiOy AlpOy(40 Å on Al) Carbon black Teflon	Substrate
**************************************	£0
103 103 103 103 103 103 103 103 103 103	y₀({10°² J m°²)

^{*}Calculated from the heat of adsorption $E_{\mu}(I|m^2)$ of the powders in solvents of different dipole mament μ . $E_{\mu} = E_{a\mu}$. E_{a} in $V|m^2$ per cm² of surface.

Girifalco, Good, Fowkes, Young equation (GGFY) permits the calculation of the interfacial energy yst.:

For LCs, γ_n^L is of the order of 10^{-2} J m⁻², %, Polyethylene has $\gamma_1 = \gamma_1^2 = 3 \times 10^{-2}$ J m⁻² thus $\gamma_{2L} = 27 \times 10^{-2}$ J m⁻².

In general the alignment of LCs on polymers is well described by the FCK rule, $^{36.77}$

II.1.6 The "smecticlike" interfacial layer

The interfacial layer between a solid and a liquid gives rise to a charge separation, which can be well described in terms of a double layer (DL) which consists of a compact ordered monolayer firmly bound to the substrate, followed by a layer of diffuse charges. This DL has not been studied in LCs where the electric charge density is not known, but its existence has been theoretically predicted and is supported by electrical measurements. It could be relevant to the higher order parameter of the interface between nematic LCs and solids or gases.

Practical LCs being ionically very pure the compact layer will be the main component.

Many experimental observations imply that the surface layer has a higher order than the bulk. Microscopic observation of nematic layers squeezed between glass slides suggested that there is a thin interfacial layer firmly bound to the glass substrate, which determines the orientation of the adjacent layer.

Infrared spectroscopy indicates that the excess order depends upon the nature of the substrate (glass, $\ln_2 O_3$, polysiloxane). ¹³⁷⁶ Additional evidence for the existence of a DL comes from observations of a film of the LC 5CB, less than 250 Å thick pulled apart in air. Such a layer has the appearance of a smectic layer with an order parameter, S_{ϕ} , 1.005 times that of the bulk value S_c . ¹³⁰

Also, whereas the surface tension of a normal liquid decreases when the temperature increases, the reverse has frequently been observed with nematic LCs. This effect could be explained by a higher surface order parameter S_{ϕ} : γ_{0L}^{0} is the surface tension of the isotropic LC, then the surface tension $\gamma_{0}^{0}(S)$ of the oriented layer is

$$\gamma^{0}(S_{\phi}) = \gamma_{0L}^{\alpha} - \alpha S_{\xi}$$
 with $\alpha \sim 2 \times 10^{-4}$ J m⁻².

Optical measurements clearly show the higher birefringence of the interfacial layer, e.g., $S_a = 0.5$, and $S_c = 0.4$ (Ref. 137) for the LC 7CB. Theoretical investigations of surface alignment have to take into account this more ordered surface layer. ¹¹ The structure of this first compact layer, which also determines the surface pressure of the adsorbed LC vapor, influences the orientation of the nematic crystal layers throughout the bulk of the LC, as does a layer of smeetic liquid crystal deposited on a substrate. ⁹⁰

The dispersive interactions of LCs with solid surfaces are strong. For metals and oxides γ_s^d is about 0.1 J m⁻² and around 3×10^{-2} J m⁻² for polymers. The dispersive component of LC surface tension has been found predominant over polar interactions $\gamma_L^d \sim 3 \times 10^{-2}$ J m⁻²; the interfacial energy is for most LCs essentially dispersive leading to the parallel alignment of the alkyl-aromatic part of the LC molecules. ¹²⁴ The polar contribution of the solid-liquid interactions will favor the parallel alignment of negative Δ_L LCs while it will lower the anchorage of positive Δ_L LCs. ¹¹³ Experimentally it is observed that cyanobiphenyls are more easily oriented homeotropically than negative esters. Silicone layers, for example, orient high positive Δ_L LCs homeotropically but not phenyl cyclohexanes.

During the initial contact between the LC and the surface, the smecticlike layer forms. Defects spreading out through the LC film are initiated by some surface irregularities. Often they evolve in time as adsorbed atmospheric components exchange with LC molecules.

The energy confined in the smecticlike layer is higher than can be added to the LC film by mechanical, electrical, or thermal action. Once it is formed, the substrate enclosing the LC film may be twisted without changing the interfa-

¹ We have not reviewed the influence of temperature on LC surface tension and alignment as our own work on the subject is insufficient on the one hand and published results too contradictory on the other hand. Some indications may be found in Refs. 3 and 111.

cial layer.34 Also its orientation at the free surface is not changed by applying a magnetic field. 138 Consequently, the physical properties of LCs are weakly affected by the very first layer (see also § II.3).

II.1.7 Nonuniform coverage

observed, depending on relative values of the LC elastic constants. 139 This situation is likely to occur in the case of absorbed surfactants at low coverage¹⁴⁰ or when hydrolyzed chloro or amino silanes form polymeric nodules on the surface, 141 or when very thin polymeric 91 or evaporated films are deposited on the substrates. 132 In these cases one is dealing with a parallelsubstrate and the alignment layer tend to induce different orientations of the LC molecules, tilted alignment will be observed. The theoretical computation of the equilibrium director orientation shows that different tilt angles will be exposed substrate and the aligning layer will influence the LC alignment. If the When an aligning layer is deposited nonuniformly on a substrate, both the orienting substrate presenting patches of homeotropic-aligning layers.

Alternating stripes of surface materials, one giving a homeotropic and the other giving parallel alignment produce a bulk average tilt angle of 45° 139 When the bend constant, K_{33} , is larger than the splay constant, K_{22} , theoretical calculations indicate that the angles will be less than 45°, while it will be greater in the inverse case.

II.1.8 Alignment by surfactant

Almost every known class of surface active agent has been observed to influence LC alignment (see § 3, Part I).

In most cases LCs are aligned homeotropically by long chain surfactants adsorbed on inorganic substrates. Because they lower surface tension, it has often been thought that the FCK rule should apply and explain the observed

pounds or a condensed (L_2) state where the surface covered by each adsorbed Superficial films of adsorbed molecules are known to exist in four states 121; the G (gaseous) state corresponds to a weak coverage. Higher coverages lead to an L (liquid) state either expanded (L $_{ ext{I}}$) with an average surface of 50 $extsf{A}^2$ (40-70 ${
m \AA}^2$) covered by each adsorbed molecule as for long chain polar commolecule is about 22 Å² (cetyl alcohol).

Dense layers create S (solid) films with an area of 20.5 Å². Some surfactants ties of superficial films of phospholipids (e.g., lecithin) have been explained as produce both the L₁ and L₂ state with an intermediate (1) phase. The properdue to their liquid crystalline organization (Figure 7).

In order to obtain monolayers of known structure, a Langmuir balance is generally used, 124 but the technique is tedious and difficult to apply in industry. The normal practice, therefore, is to adsorb layers from solution and relate

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heir structure to that of the Langmuir monolayer. The retraction method has been shown to form good monolayers 142 although they present slight differences in their properties compared to the former, 143 probably because of the incorporation of solvant into the layer.

of the curves and homeotropically in the lower one. "On the completion of the adsorbed monolayer on the surface of a solid, the orientation of hydrocarbon nonpolar molecules on a polar solid stand upright as shown in 5b. However, if The investigation of the structure of layers adsorbed from solutions is still in progress¹⁴⁴ and their are few cases where it is complete. In Figure 4 we have sults are assumed to indicate the influence of the carbon content of the chain on the substrate interfacial energy, the relation of LC alignment is not so clear, as in every case MBBA is aligned parallel to the substrate in the upper portion plenty of area is available, as in a dilute film, the polar-nonpolar molecules type of orientation. If the vapor becomes saturated and the liquid of the film gathered experimental values of the critical surface tension of glass covered with adsorbed layers of fatty derivatives of increasing chain length. If the remolecules is parallel to the substrate as shown in Figure 5, whereas polaralso lie down. As the film thickens and becomes multimolecular, the orientation in the first layer of polar-nonpolar molecules preserves the perpendicular wets the solid, the film is relatively thick (about seven molecules thick for water at 25°C) and the molecules in the surface of the film have approximately the same orientation as in the surface of the liquid itself". 124

In addition, the packing density of fatty acids depends on their chain ength. 142,145 This could be an explanation for the tilted alignment of MBBA on surfaces covered by retracted layers of C10-C6 fatty derivatives.

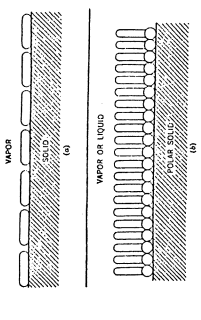


FIGURE 5 Orientation of molecules in complete monolayers. (a) Hydrocarbon molecules. (b) Polar molecules on the surface of a polar substrate. (From Ref. 124).

on these layers takes the surfactant orientation (Figure 6). surfactant molecules orient perpendicular to the substrate." MBBA deposited parallel to the substrate at low concentration while at higher concentration the tion isotherms. Cetyl ammonium bromide (CTAB) solutions in water adsorb Liquid crystal alignment may be understood from the surfactant's adsorp-

orientation whereas on layers of lower packing density homeotropic orientalayer structure.117 On tightly packed layers, cyanobiphenyls show a tilted cules are imbricated in the surfactant layer. 148 tion is observed. Experimental evidence leads to the conclusion that LC mole-7) depending on their surface pressure, retraction on glass substrate keeps the Phospholipids (e.g., lecithin) monolayers have different structures (Figure

structure: cyanobiphenyls are much more easily aligned homeotropically than phenylcyclohexanes. The efficiency of a surfactant to align LCs also depends on its molecular

derivatives. more permanent effects to result from the adsorption of long chain silane As silanes may form covalent bonds with oxide substrates one would expect

nodules very often form on the surface¹⁴, but can be avoided by anhydrous conditions. ¹⁴⁹ Very little is known about the structure of silane monolayers. Polymeric

alignment (Table V), is also known to be densely adsorbed on tin oxide from a OTS. 3-(2-Aminoethyl)aminopropyl TMS, used to facilitate homogeneous All our test LCs aligned homeotropically on oxide substrates covered with corresponds to close packing of Si-O bonds associated in three dimensions. 150 1.5 × 10¹⁴ mole/cm², as compared to 5 × 10¹⁴ for ¹⁴C arachidic acid, which forms dense monolayers.14 Carbon-14 counting shows a surface density of 5% solution in toluene. 149 Octadecyltrichlorosilane (OTS) adsorbed from nonaqueous solutions

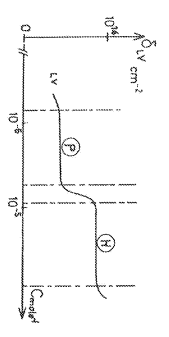


FIGURE 6. Variation of the superficial density (850) of adsorbed cetyl trimentyl ammonium bromide (CIAB) with its concentration in solution. (From Ref. 157).

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GASEOUS (G) STATE

LIQUID (L) STATE

smectic lamella

nematic lamella

CITOS (S) STATE

FIGURE 7 Schematic representation of phospholipid (e.g., lecihin) monolayers. (From Ref.

23

Surface hydrolysis 23.11

Liquid esters have surface tensions of $28-30 \times 10^{-3} \, \mathrm{J} \, \mathrm{m}^{-2}$ and should spread metals they do not always do so on glasses or oxídes. This irreproducibility has on high energy surfaces such as metals and oxides. Whereas they do spread on been related to the ester hydrolysis catalyzed by adsorbed water, it the ester being unable to spread on an adsorbed layer of its own hydrolysis products.

The same is believed to be true of the occasionally observed homeotropic alignment of LC esters of Schiff bases (SB) on glasses and some oxides (Al2O3, In2O1). This effect is normally only observed if the surfaces have previously been washed in acid. This alignment disappears when the substrate is fired and is not observed with pure LCs of other molecular structure. On dry substrates, LC esters, as opposed to liquid esters, 143 are flat on the surface due to the carboxylic group interacting strongly with the oxide. 151

tions higher than could be adsorbed from solutions; 1% concentrations of main stuck to the substrate even after a heating cycle or the emptying and refill-ing of a cell. A related impurity induced homeotropic aligning effect is duce alkylbenzoic acid or alkylamine, respectively, both of which are known these substances in the LC are required to induce the same alignment. An indi-Photooxidation of the LC forms oxyderivatives 435 that induce homeotropic SBs are completely hydrolyzed during HPLC on silica columns20 and are known to be very sensitive to moisture. 182 Hydrolysis of ester or SB LCs proto induce homeotropic alignment. Surface hydrolysis leads to local concentracation that surface hydrolysis has occurred is the formation of defects that reobserved when LC layers are either left in air or submitted to light irradiation. alignment. In the case of unsealed cells the perpendicular orientation proceeds from the outer edge.

II.1.10 Measurement of LC surface tension

cations of the choice of one method. Capillary, do Nouy, Wilhemy methods This section is not devoted to a discussion of experimental surface tension measurements that would be found, for instance, in Ref. 121, but to the impliimply the contact of LCs with surfaces,

derstand the difficulties, generally underestimated, of obtaining reliable From the preceding considerations of the LC-solid interaction one can unvalues; after long and careful studies it has been concluded that surface tension of MBBA is "impossible to measure". In

As LCs lie generally parallel to solid surfaces the above quoted method should give a value near that of yk.

to the free surface the value of γ_1 would be obtainable with a precision of about 5%. In fact, even in this method the solid-liquid interactions are predominant. Only in the hanging drop method are the solid-LC interactions minimized and the yea values should be obtained. 116 As LCs often orient perpendicular

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The wide range of literature values may correspond to different aligning conditions and indicate an anisotropy $\Delta\gamma$ of about 10 erg, in agreement with theoretical considerations.

11.1.11 Summary of Section II.1

stead, the experimental yevalue, which gives a rough expectation of the direction of LC alignment. This can only be exact when the ratio of the polar and As the solid surface tension, y, is not measurable, one has sought to use, indispersive components of both the solid and the LC are equal.

A quantitative description of a LCs alignment requires knowledge of the polar and dispersive parts of surface tension as polar interactions depend on a LCs dielectric anisotropy and may be predominant.

ity of recent work suggests the validity of the FCK rule; much more work has to be done to get reliable values of a LCs surface tension. As long as experiyscs betwen 30 and 40, critical evaluation of the FCK rule, which requires a Determination of a LCs surface tension is, currently, not accurate enough to allow a precise comparison of substrate and LC surface tension. The majormental values of ymaga continue to be found between 38 and 28 erg/cm², or knowledge of y to better than I erg/cm?, is meaningless.

In our opinion there are no sound experimental data that invalidate this rule. Discrepancies indicate experimental error and we will suggest its use as a air estimate of the expected alignment of a LC.

il.2 Elastic ortentation on grooved surfaces

II.2.1 Geometrical factors

On high energy substrates ($\gamma_{\rm SV} \ge 4 \times 10^{-3} \ {\rm J \, m^{-2}}$), LCs align parallel to the substrate. Sometimes, the flow of a LC drop will produce a nearly uniform alignment, albeit with many defects. This orientation occurs most easily when the LC is in its isotropic state. Very slow cooling favors uniformity and good alignment is obtained on cooling in a magnetic field.44

tional polishing, tangential evaporation, tangential ion-beam etching of These methods are impractical and uniformity is more easily obtained on grooved surfaces. Grooves are produced by rubbing (glass plates), unidirecoxides, or the formation of a grating on the substrate, 47,54

aematic interactions at the shear plane; the LCs anchorage is fairly weak and lations of the clastic energy added to the LC bulk by a periodic surface deformation show⁴³ that the LC layer has an energy which is greater when the LC Once the first interfacial layer is formed, the LC layer aligns through the defines a slippery surface. (see § II.3.1.) In these conditions, theoretical calcudirector is perpendicular to the groove direction d (but parallel to the substrate) than when the director lies in the direction of the groove.

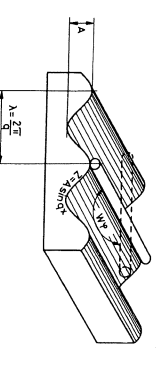


FIGURE 8 The sinusoidal model of a grooved surface. The LC molecules have higher clastic energy in the orientation perpendicular to the plane than parallel to it.

If the surface profile (Figure 8) is defined by the depth, A, of the grooves and their spacing $\lambda=2\pi/q$

$$Z = A \sin qx \qquad U = \frac{K}{4} A^2 q^3$$

where K is the LC elastic constant. Table IX summarizes the profiles obtained by different grooving techniques and the related increases in energy for alignment parallel to the grooves. Photolithographically formed gratings allow the characterization of the periodic profile that influences the alignment. For 250 Å deep grooves uniform alignment of MBBA is obtained for

$$\lambda < 3.4\mu$$
.

In general the ratio A/λ is smaller than 1.

TABLE IX

Estimated values of depth and spatial distribution of grooves on various substrates in the sinusoidal approximation

	Evaporated SiO _x 60° Gratings	Diamond pencil scratches	Rubbed glass Polished glass (diamond no	Surf. preparation
	SiO, 60°	oencil	ass lass	paration
***************************************	100 250	10 000	10 200	Depth A (Å)
	200–400 3200	<10 µ	200 1000	Spacing (Å)
	6 × 10 ⁻⁴ 1.2 × 10 ⁻⁴	15 × 10 ⁻⁷	8 × 10 ⁻⁵ 2.4 × 10 ⁻⁴	Energy U J m ⁻²
	44(a) 54	47	43(a) 43(b)	Ref.

^{*}U is the excess energy of the nematic liquid crystal aligned perpendicular to the grooves over its energy when it is aligned parallel to them. Calculated from $U=(1/4)KA^2q^3$ assuming a unique elastic constant $K=10^{-11}$ N.

If the different elastic constants of the LC are taken into account, calculations⁴⁷ indicate that for $K_{22} > 3K_{33}$ a metastable configuration exists where n is \perp to d which cannot be observed, as $K_{22} \sim K_{33}/2$ for LCs.

Grooving does not influence homeotropic alignment provided the surface profile is symmetrical (Figure 9a'). Silanes or silicones deposited on grooved surfaces still induce homeotropic alignment. ^{20,157} Sawtooth profiles may produce an average tilted orientation in the bulk of a LC⁴⁴ (Figure 10), as will an unsymmetrical coating of the grooves. ¹⁰⁸ Considering that the LC molecules are either lying parallel to the substrate or orthogonal to it ^{111b} three configurations of the LC may exist. ⁷ In the case of Figure 9a' the elastic energy of the LC layer is not a minimum and it will not be observed on a sinusoidal profile as described in Figure 8.

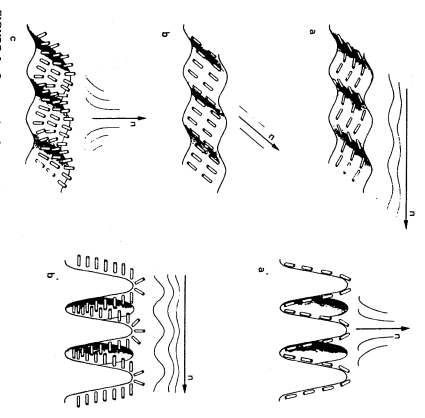


FIGURE 9 On grooved surface, parallel or homeotropic alignment may be observed depending upon the spatial period to amplitude ratio.

J. COGNARD

A three dimensional profile will create an alternation of peaks and valleys and constrain the LC to take the configuration of Figure 9b.

mension the following situation may occur: if the substrate tension energy is high the molecules will lie flat on the surface but the bulk average will be homeotropic (similar considerations are found in Ref. 57). Lowering the sur-Depending on the width to depth ratio as compared to the molecular diface tension energy by surfactant adsorption will orient the molecules perpendicular to the surface resulting in an homogeneous alignment.

at least, three molecular dimensions (i.e., 60 Å as LC molecules typically have Experimental evidence will be published later.‡ It requires that the depth of the hole be bigger than its width and the spacing of two pits is of the order of, length of 20 Å and width of 6 Å).

II.2.2 Mixed alignment

The alignment of LCs on parallel orienting surfaces with patches of a homeotropic aligning agent has been considered in § I.1.7.

osition of a dense layer of a homeotropic aligning agent (see § II.1.8.) on a ented normal to the surface, a distortion exists in the LC layer. When such a homeotropic aligning layer is deposited on a nonsymmetric layer, producing a tilt angle ϕ , the complementary, $\phi + \pi/2$, tilt angle is observed (Figure 3). 5.20.157 +Obliquely ($\theta > 15^{\circ}$) evaporated SiO_x layers align LCs homogeneously. An additional tangential evaporation ($\theta < 15^{\circ}$), which would otherwise produce a tilt angle of 25°, permits the selection of any orientation (土3°) of the LC molecules between 0° and 25°, as the asymmetry of the grooves increases. Depnot change the homeotropic orientation, 69,17 but as the first monolayer is orisymmetrically grooved surface (either rubbed or obliquely evaporated) will

II.3 Anchoring energies

11.3.1 Anchoring energies and the interfacial layer

The interactions of the LC molecules with the surface have to be taken into account when considering the LCs static deformation.

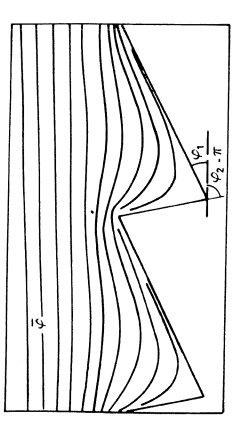
The energy of interaction, W_{\bullet} , can be expressed phenomenologically as the sum of an isotropic energy term, Wo, an in-plane (twist or torsion) anchorage and an out-of-plane (azimuthal) anchorage:

$$W_{\bullet} = W_0 + W_{\phi} + W_{\theta},$$
 (3.1)

where the angles are as indicated on Figure 11. The angular dependence of $W_{m{\phi}}$ and We can be described153 by

ALIGNMENT OF LIQUID CRYSTALS

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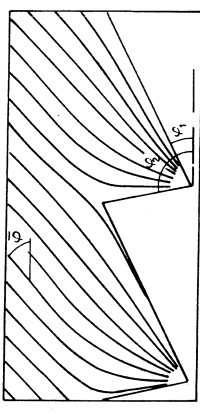


FIGURE 10 Unsymmetrical grooves may induce tilted alignment (case b) as evidenced by a rheographic model. (From Ref. 44). (a) $\phi = \phi_1$ on surface 1, $\phi = \phi_2 - \pi$ on surface 2, $\overline{\phi} = 0$. (b) $\phi = \phi_1$ on surface 1, $\phi = \phi_2$ on surface 2, $\overline{\phi} = \phi_1[\pi/(\pi - \phi_2 + \phi_1)]$.

$$W_{\phi} = B_{\phi} \sin^2 \phi, \tag{3.2}$$

$$W_{\theta} = W_0 + B_{\theta} \sin^2 \theta. \tag{3.3}$$

In the physical chemistry of surfaces, interfacial interactions are usually described by the surface tension of both phases.

When a liquid of surface tension year comes into contact with a solid of surface tension ysv°, and interfacial layer builds up whose energy ysL (identical to

$$\gamma_{\rm SL} = \gamma_{\rm SV} + \gamma_{\rm LV} - W_{\rm a}, \tag{3.4}$$

[†] Similar observations have been reported in Appl. Phys. Lett. 36, 144 (1980) by W. R. Heffner

[‡] F. Gharadjedaghi, "Liquid crystal alignment techniques producing controllable tilt angles" presented at the 4th Liquid Crystal Conference of Socialist Countries, October, 1981.

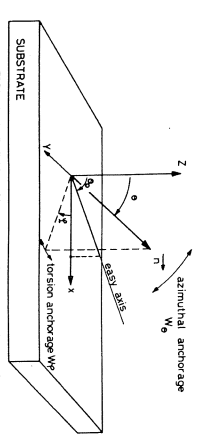


FIGURE 11 Definition of in-plane and azimuthal anchoring energies

where γ_{SV^0} is the solid surface tension at equilibrium with liquid saturated vapor (which is related to the solid-vacuum interface free energy γ_S and the adsorbed liquid layer surface pressure π_0 by $\gamma_{SV^0} = \gamma_S - \pi^0$) and W_a is the adhesion energy which may be approximated¹⁵⁹ by

$$W_{\bullet} = 2\sqrt{\gamma_{\text{SV}}\gamma_{\text{LV}}}.\tag{3.5}$$

The solid surface energy, γ_8 , is not directly measurable and depends on the surface preparation and history, but for inorganic substrates γ is calculated to be >0.2 J m⁻², in agreement with values obtained from heat of immersion measurements. For polar liquids the surface pressure π_0 exceeds 0.1 J m⁻² leading to $\gamma_8 v^6$ values >7 × 10⁻² J m⁻² (Ref. 117) (the surface tension of an inorganic substrate covered with a water layer).

In the case of a liquid of $\gamma_{LV} = 3 \times 10^{-2} \, \text{J m}^{-2}$, which is typical for LCs, the interfacial layer energy γ_{SL} is obtained from (3.4) and (3.5) giving $\gamma_{SL} = 8 \times 10^{-3} \, \text{J m}^{-2}$.

The thickness of the interfacial layer is not known precisely but it extends over at least 100 Å in a resistive liquid with LC molecular dimensions.

From the interfacial energy profile as sketched in Figure 12, one sees that the anchoring energy varies with the distance from the surface.

The excess energy of the interfacial layer produces in nematic LCs a higher ordering which creates the smecticlike layer described in § II.1.6. As surface tension exerts a force normal to the substrate, there is no dependence of the interfacial energy on the in-plane orientation of the molecules.

II.3.2 In-plane anchorage energy

The torsion anchorage, W_{ϕ} , results from substrate nonuniformity which may be controlled, for instance, by uniform grooving of the surface (Figure 8), in

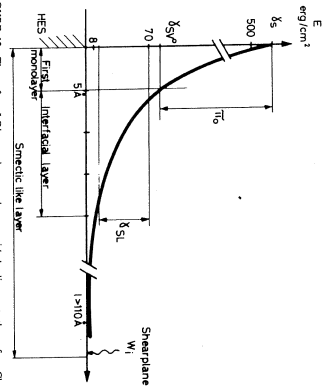


FIGURE 12 The surface-LC interaction energies vary with the distance to the surface. Close to the surface an ordered layer is related to the smectic organization. There exists a shearing plane at a distance where the energy of elastic deformation equals the excess energy due to surface interactions.

which case W_{ϕ} may be calculated (see § II.2; $W_{\phi} \equiv U$). W_{ϕ} may be roughly estimated from the width, h, of surface disclination lines¹⁵³

$$W_{\phi} = \frac{\pi^2 \, Kh}{2 \, d^2}$$

(where d is the LC layer thickness), or measured from the variation in twist angle of a twisted layer subject to a magnetic field. ¹⁵⁵ As both experimental measurements and the elastic alignment theory relate to the alignment beyond the smectic layer, calculated and experimental values, as compared in Table X, show that the orders of magnitude are in fair agreement. In every case the torsion anchoring energy is higher than the bulk twist energy

$$W_{\rm twist} = K \, \phi_{\rm r}^2 / 2d \approx 10^{-6} \, {\rm J \ m^{-2}}$$

 $(\phi_T = \text{twist angle, See Ref. 158}).$

In a twisted nematic layer, minimizing the bulk free energy leads to the equilbrium condition $d\phi/dz = \text{constant}$. In the vicinity of the substrate it is

	We exptl (J m ⁻²)	⁸ (^{5−} m L) olso _{&} W	Substrate
J J	2 × 10.4 (153)		SealO
COGNARD	(\$\$1) ² 01 × 7 (\$\$1)	8 × 10 ₋₂	Rubbed glass
n	10-3	►01 × S	%SiOx evap 30°
	(5) ₅₋ 01 × 5 ~	•••	Heptylamine on glass
	(ç) _← 01 × ç ≥		Trimethyltrichlorosilane on glass

• $W_{\phi} \equiv U(T_{able} IX) = 1/4 \text{ K A}^2 \phi^3 (1 \text{ erg/cm}^2 = 10^{-3} \text{ J m}^{-2}).$

Wall Surface Bulk nematic region and the "extra polation length." Weak in-plane anchoring

FIGURE 13 Definition of the "extrapolation length." Weak in-plane anchoring corresponds to $b \leqslant a$. (From Ref. 156).

expected that this condition does not hold¹⁵⁶ and extrapolation of the linear variation of $\phi_{(2)}$, to $\phi = 0$ occurs at z = -b (Figure 13). The "extrapolation length" $b = K/B\phi$ would in this case define the condition of strong anchoring. The twisted nematic layer behaves as if it had a thickness d + 2b; thus the anchoring is strong if $2b \leqslant d$.

The threshold voltage under applied electric field is given 157 by

$$V_{\rm th} = \frac{\pi}{d+2b} \sqrt{\frac{4\pi K}{\Delta \epsilon}} \left(1 - \frac{2K}{B_{\phi}d} \right). \tag{3.6}$$

As the torsion anchoring energy on grooved surfaces may vary from 5×10^{-6} to 10^{-4} J m⁻² (Table IX) the extrapolation length may reach 2μ for a typical LC cell ($d = 10^{-5}$ m) ($K = 10^{-11}$ N). In fact we have not found experimental evidence for the extrpolation length.

II.3.3 Azimuthal anchorage

As W_{ϕ} is independent of the interfacial interactions on uniform surfaces, the anchoring energy reverts to the energy of the azimuthal anchorage, which is equal to the difference between the interfacial energy when the LC is aligned parallel or perpendicular. Let $\xi(0)$ and $\xi(\pi/2)$ be the values of the physical property ξ in both orientations

$$= W_0 + B_\theta \sin^2 \theta = \gamma_{\rm SL}(0) + \left[\gamma_{\rm SL} \left(\frac{\pi}{2} \right) - \gamma_{\rm SL}(0) \sin^2 \theta \right]. \quad (3.7)$$

In Eqs. (3.4) and (3.5), π_0 , $\gamma_{\rm SL}$ and $W_{\rm a}$ are θ dependent, and

$$B_{\theta} = \gamma_{\mathrm{SL}} \left(\frac{\pi}{2} \right) - \gamma_{\mathrm{SL}(0)}$$

$$= \left[-(0) - \left(\frac{\pi}{2} \right) \right] \cdot \left[-(\pi) - (\pi) - (\pi) - (\pi) \right] \cdot \left[-(\pi) - (\pi) - (\pi) - (\pi) \right] \cdot \left[-(\pi) - (\pi) - (\pi) - (\pi) - (\pi) \right] \cdot \left[-(\pi) - (\pi) - (\pi) - (\pi) - (\pi) - (\pi) \right] \cdot \left[-(\pi) - (\pi) - (\pi) - (\pi) - (\pi) - (\pi) - (\pi) \right] \cdot \left[-(\pi) - (\pi) - ($$

glass/MBBA interface to be 6.2×19^{-3} J m⁻² (8 c) and is calculated to be pected to be large. The LC surface tension anisotropy, $\gamma = \gamma_{\rm LV(0)} - \gamma_{\rm LV(\pi/2)}$ is estimated to be $+4.5 \times 10^{-5}$ J m⁻² for MBBA⁶⁶ and 5×10^{-6} J m⁻² for 5 CB pendicular orientation has been deduced from experimental data at the (102 a). The difference in adhesion energy $W_{\mathbf{a}}$ between the parallel and perthe surface pressure in the literature. As on inorganic substrates LCs prefer to be parallel to the substrate, $\pi_0(0) > \pi_0(\pi/2)$ and the anisotropy is not ex-There is no reliable data on these values for LCs. No account has been taken of $= \left[\pi_0(0) - \pi_0\left(\frac{\pi}{2}\right) \right] + \left[\gamma_{LV}\left(\frac{\pi}{2}\right) \gamma_{LV}(0) \right] + \left[W_{a}(0) - W_{a}\left(\frac{V}{2}\right) \right]. \quad (3.8)$

energy. From (3.8) one gets: Thus the energy in the interfacial layer is mainly determined by the adhesion

 7.16×10^{-3} J m⁻² for PAA on glass (102).

$$B_{\theta} = \gamma_{\text{SL}(\pi/2)} - \gamma_{\text{SL}(0)} \sim W_{\text{a}}(0) - W_{\text{a}}\left(\frac{\pi}{2}\right) = 6 \times 10^{-3} \text{ J m}^{-2}.$$

dary conditions to be satisfied at the surface. interfacial layer cannot be modified by elastic deformation of the LC where the energy is at most 10⁻⁵ J m⁻². In this case, ¹³⁸ one should not minimize the bulk plus surface energy, but minimize the bulk energy and require the boun-This large difference in the surface energy means that the orientation of the

Writing the LC layer energy as

$$F = \frac{1}{2} W_{s} \theta^{2}_{(0)} + \int_{0}^{d} \frac{1}{2} K \left(\frac{d\theta}{dz} \right)^{2} dz$$

the substrate will change its orientation beyond a distance d_c : homeotropically oriented and subject to a magnetic field directed parallel to the flea to lower the weight of the combined system. A LC layer of thickness d, is tantamount to putting a flea on an elephant and minimizing the weight of

$$d_{\rm c} = \frac{3K}{2(\gamma_{\perp} - \gamma_{\parallel})} \tag{3}$$

substrate. Azimuthal anchoring energies are either estimated from surface drilled in a crystal sheet, and subjected to a magnetic field perpendicular to the as observed 138 on a homogeneously aligned drop of PAA deposited in a hole disclination lines 153 or measured by magnetic deformation of homeotropically

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TABLE XI

Experimental values of the angular coefficient B_{θ} of $(W_{\theta} = W_0 + B_{\theta} \sin^2 \theta)$ azimuthal anchoring energy W_{θ}

i											
	in2O3 + NH2C16H33	In ₂ O ₃ + lecithin	In ₂ O ₃ + DMOAP	Glass + DMOAP	Glass + lecithin	Class + NH ₂ -C ₇ H ₁₅	Class + NH ₂ C _n H _{2n+1}	Class + R,N Mc3X	SIO _x (30° + 5°)	Air	Substrate
	MBBA	MBBA	MBBA	MBBA	MBBA	MBBA	MBBA	MBBA	6CB	5CB	TC
	3.5	6	10	6	w	2.6	2	w	2.1	4	$(10^{-6} \text{J m}^{-2})$
	MD	ΜD	MD	M	MD	SD	MD	M	Δn	DA	Method*
	75	75	75	75	75	5(a)	155	75	109	120(a)	Ref.

^{*}Method: DA, differential anchoring: Δn , birefringerence; MD, magnetic deformation; SD, surface disclination. The order of magnitudes of B_{θ} compared to the experimental surface tension anisotropy. MBBA: $\gamma_{\parallel} - \gamma_{\perp} \sim 4.5 \times 10^{-5} \text{ J m}^{-2}$ [Ref. 75(b)]; 5CB: $\gamma_{\parallel} - \gamma_{\perp} \sim 5 \times 10^{-6} \text{ J m}^{-2}$

 $B_{\theta} \sim \gamma(0) - \gamma(\pi/2)$. tion of the LC layer where the LC is not affected by the surface potential and between calculated and experimental values would correspond to a deformacomparable to the LC surface tension anisotropy (Table XI). This difference oriented layers where (3.9) should apply. The B_{θ} values obtained are weak and

strate, even though the anchorage is strong, weak interactions are observed as layer. The same situation is probably observed on most adsorbed surfactant layers. With $B_{\theta} \approx 5 \times 10^{-6}$ J m⁻² the anchorage parameter of Ref. 158 the perturbed part of the nematic layer interacts with its smecticlike interfacial On inorganic layers where a LC would normally align parallel to the sub-

$$\lambda = \pi \frac{K}{B_{\theta}d}$$

is about 1 for a LC with $K \approx 10^{-11}$ N and a cell thickness of $d = 10\mu$

strongly reduced and the interfacial layer may disappear, but very often hochorage than now known. On polymeric films surface interactions are meotropic alignment is observed on these surfaces. Thus one may wonder whether it would be possible to observe weaker an-

II.1.4 Conclusions

As phrophesized sixty years ago²³;

"Les actions auxquelles obéissent les liquides anisotropes, qui se révèlent par des structures si variées, pourront sans doute plus tard se résumer en quelques lois très simples.

servations très nombreuses recueillies jusqu'à ce jour, qu'elles feront jouer un rôle très ieures qui fixent l'orientation du liquide par rapport à lui-même dans toute l'étude de Bien que ces dernières ne soient pas connues, on est en droit de penser, d'après les obimportant aux actions superficielles qui règlent l'orientation du liquide au voisinage de la surface, dans la couche capillaire, et les distingueront nettement des actions intérsa masse. Aux premières se rattachent par exemple les plages orientées à la surface du verre: aux secondes les coniques focales, les fils."

of the molecules minimum energy, LC molecules open wide fields to Constituted of an aromatic core having strong dispersive interactions with the substrate, an alkyl chain tending to orient perpendicularly to the surface as the strates, pure LC and well defined aligning layers free of solvents and ionic conchain length increases, and a polar group which may counteract the other part Mastering the alignment of LCs requires the control of the smecticlike interfacial layer. In addition to the difficulties of obtaining uniform and clean subtamination, LC molecular structure gives rise to complicated interactions. speculation.

if all aspects of LC orientation are not under control, as is also the case for the Theoretical expectations are summarized in Table (XII). Nevertheless, even physical chemistry of surfaces, reproducible and well defined alignment can be obtained, as described in the last paragraph.

action with the surface as in suspended films, on low energy polymers and on In general, inorganic substrates lead to nonuniform parallel alignment, the polymeric layers. Homeotropic alignment is observed in the absence of interconditions that lead to homeotropically aligned LCs on these surfaces have not been defined. Rubbing will provide uniformity; it is best achieved on soft adsorbed surfactant layers. The very first organized layer reaches its equilibrium slowly and determines the final nematic crystal structure.

III RECOMMENDED PROCEDURES

and temperatures vary. It is therefore desirable to use procedures which have ing a LCs physical properties to be performed without the use of cumbersome Alignment problems arise frequently in laboratory experiments where LCs been well tested. The use of good alignment layers allows experiments involvThe success of any aligning process depends upon the substrate and the cleaning procedures employed. The alignment of a LC may be studied on metal or crystal surfaces, but it is more usual to employ transparent substrates.

should be aware of the many varieties of glass compositions available and the various manufacturing methods (pulled, floated, fire polished) which lead to Glass, quartz, silicon coated glass, or transparent conducting coatings (SnO₂, In₂O₃)—patterned or otherwise—are most frequently employed. One

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TABLE XII

Alignment expected from substrate/LC interaction characterized by the dispersive and polar components of the solid and LC surface tension

**	Δε < 0	No cases	polar surfaces $(\gamma_{\bullet}^{p} \gg \gamma_{\bullet}^{d})$	$\gamma_{ullet} > \gamma_{ m d}$			₽	⊣	>-	=
γe > γ.	Δε>0			have			Γq	·	- 4-	
به:	Δε < 0	×		=		4	×	=	\	٦
, , ,	Δε > 0*	٩Τ		⊣		The state of the s	×	=	¥	٦ ا
crystal	Surf. state	Smooth		Sym grooves	Unsym grooves	Tridimensional ^e pit array	Smooth	Sym grooves	Unsym grooves	Tridimensional ^e pit array
Liquid crystal Surface	Surf. tension $\gamma_\bullet = \gamma_\bullet^P + \gamma_\bullet^d$		p * A α * Λ	Activated oxides diacids			*	Humid oxides polymers fatty deriv.		

^{*} Δε gives an indication of the LC polarity

^bThis case is not experimentally very well documented.

Regularly repeating tridimentional array of peaks and valleys.

d Homeotropy due to tridimensional array does not depend upon LCs nature while that due to absence of interaction does.

^{· +} and + = tilted alignment.

different interactions with the LC to be aligned. The use of an acid bath to clean the substrate changes its properties towards certain classes of LCs. These considerations are not too important if the substrate is to be coated with a polymeric or oxide layer, but they are essential when silanes or surfactants are to be employed as aligning agents. More reproducible results are always obtained on an underlying coating.

The following methods have been widely used with success and give good results with most combinations of substrate and LC.

III.1 Cleaning

Cleaning of solid substrates is still an art despite the considerable amount of work done on it. ¹⁶⁰ Our studies use either a detergent bath or dipping in boiling sulfuric acid followed by thorough 18 M Ω water rinse and centrifuge drying.

Other methods may be found in the literature, as for instance in Ref. 159. A very important step in any cleaning procedure is the drying, which has to be effected rapidly, i.e., under a high pressure stream of gas.

III.2 Parallel alignment

Alignment by the rubbing of a substrate has been reviewed. In the laboratory, uniform alignment will be difficult to obtain reproducibly on every substrate. Industrial processing employs sophisticated machines. Rubbing by hand with a woven material or a polishing paste can only be expected to give uniform and reproducible results on soft substrates (glass or polymers).

III.2.1 Rubbed PVA

Thin polymeric films are best deposited by spin coating. A spinner may easily be constructed for laboratory uses. Polyvinyl alcohol (PVA) is commercially available in different grades (Wacker Polyviol®), but it appears that its aligning properties are not related to the grade used. We currently use the M 13/140 grade, which is very soluble and produces uniform films.

An aqueous solution (3%) is filtered on 0.5μ millipore filters and deposited on a spinning (4000 rpm) substrate at room temperature. The deposited film $(d \sim 1000 \text{ Å})$ is dried at 80°C for 30 min and subsequently rubbed with a textile cloth. Uniform alignment with a very low tilt angle is obtained on all substrates and with most LCs.

III.2.2 Rubbed polyimide

Although PVA forms good aligning layers it is water and temperature sensitive. Polyimide is a very stable polymer. Films may be formed on a solid substrate from a monomer solution of an acid anhydride and a polyamine. Such solutions in N-methylpyrrolidone are commercially available (Rhône Pou-

lenc, Nolimide 32). The deposition of a uniform film is more difficult than with PVA.

Spin coating is effected as described above from a 1/10 diluted solution and the resulting deposit dried at 80°C for 30 min. The soft prepolymer is then rubbed. Polymerization is achieved by curing in two steps, 130°C, 30 min, followed by 200°C, 30 min.

III.2.3 SiOx evaporation

If an evaporator is available, SiO_x evaporation is the easiest way to obtain stable, uniform, reproducible alignment in the laboratory, provided that the substrate is flat and homogeneous. Uniform alignment with varying tilt angles may be obtained by variation of the angle between the substrate plane and the direction of the incident beam. Parallel alignment is obtain for a 30° angle.

Pure silicon monoxide (Balzers 99.99) is evaporated by techniques which are well known 161; boat temperature 1300–1400°C, $p \approx 10^{-4}$ mm Hg, air bleeding. Typical thicknesses are 200–300 Å. (MgF₂ may also be used.)

III.3 Homeotropic alignment

There are at present no entirely satisfactory methods to obtain easy, reproducible homeotropic alignment free of complications (short lifetime, conductivity increase, etc.); for experimental evaluation the following methods give good results.

III.3.1 Lecithin (DMOAP)

The easiest way to obtain a homeotropically aligned LC layer is to sandwich it between lecithin coated layers. A solution of natural egg lecithin (Merck 5331) in a volatile solvant (e.g., alcohol) is used and the substrates are either spin-coated or dipped in the solution. The concentration is unimportant but the use of a dilute (1%) solution avoids the formation of spots. The coating is rinsed with the solvent and dried at 80°C for 30 min.

Alignment of most LCs is obtained in a direction orthogonal to the substrate plane without any tilt. There is no variation of the orientation with temperature, but the LC conductivity is increased. With the same technique DMOAP [dimethyloctadecyl-3-(trimethoxysilyl)propylammonium chloride, Petrarch 09745] gives similar results and drawbacks but is somewhat more substrate sensitive.

III.3.2 DMCS

For the alignment of positive LCs (the technique fails in the case of many negative $\Delta \epsilon$ LCs) a polymethylsiloxane, insulating, homeotropically aligning layer may easily be formed on the substrate from a Dimethyldichlorosilane

3

(Fluka 40140) solution 10% in toluene) by dipping the plate for 15 min at room temperature. The coating is ninsed with isopropyl alcohol and cured at 180°C for 30 min.

iii.4 Tilled alignment

Tangentially evaporated SiOx

face state and the LCs elastic constants. It is thus difficult to know with precision the tilt angle one will obtain with different LCs sandwiched between the As previously stressed, tilt angle is a bulk average, depending on both the sursame plates. A tilt angle of 15° to 30° will be obtained on tangentially evapoated SiOx. The evaporating conditions are the same as in § III.2.3.

Crossed evaporation 111.4.2

angentially evaporated SiO, on top of a 30° evaporated layer (fow angles) or the reverse (high angles). Care must be taken to rotate the substrate by 90° between each evaporation as the nematic director is oriented perpendicular to To obtain angles between 0° and 30° it is possible to deposit a thin layer of the evaporation direction for tangentially evaporated layers.

"Titted homeotropic" alignment III.4.3

To obtain a slight tilt angle from the direction normal to the substrate a homeotropic aligning agent (e.g., Lecithin or DMOAP) is combined with one of the above methods (4.1., 4.2.). The procedure is as outlined in § III.3.

Acknowledgments

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